

This Page Is Inserted by IFW Operations
and is not a part of the Official Record

BEST AVAILABLE IMAGES

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images may include (but are not limited to):

- BLACK BORDERS
- TEXT CUT OFF AT TOP, BOTTOM OR SIDES
- FADED TEXT
- ILLEGIBLE TEXT
- SKEWED/SLANTED IMAGES
- COLORED PHOTOS
- BLACK OR VERY BLACK AND WHITE DARK PHOTOS
- GRAY SCALE DOCUMENTS

IMAGES ARE BEST AVAILABLE COPY.

**As rescanning documents *will not* correct images,
please do not report the images to the
Image Problem Mailbox.**



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification: C12N 15/13, A61K 39/395, C07K 16/00, C12N 5/10, C12N 15/63	A1	(11) International Publication Number: WO 00/29584 (43) International Publication Date: 25 May 2000 (25.05.2000)
(21) International Application Number: PCT/US99/27153 (22) International Filing Date: 16 November 1999 (16.11.1999) (30) Priority Data: 60/108,945 18 November 1998 (18.11.1998) US (60) Parent Application or Grant GENENTECH, INC. [/]; (). CHEN, Yvonne, M. [/]; (). LOWMAN, Henry, B. [/]; (). MULLER, Yves [/]; (). WENDY, M., Lee ; ().		Published
(54) Title: ANTIBODY VARIANTS WITH HICHER BINDING AFFINITY COMPARED TO PARENT ANTIBODIES (54) Titre: VARIANTS D'ANTICORPS A AFFINITE DE LIAISON SUPERIEURE PAR RAPPORT AUX ANTICORPS D'ORIGINE		
(57) Abstract		
Antibody variants of parent antibodies are disclosed which have one or more amino acids inserted in a hypervariable region of the parent antibody and a binding affinity for a target antigen which is at least about two fold stronger than the binding affinity of the parent antibody for the antigen.		
(57) Abrégé L'invention concerne des variants d'anticorps par rapport à des anticorps d'origine, caractérisés en ce qu'un ou plusieurs acides aminés sont insérés dans une région hypervariable de l'anticorps d'origine, et présentant une affinité de liaison vis-à-vis d'un antigène cible au moins deux fois plus forte que l'affinité de liaison de l'anticorps d'origine pour l'antigène considéré.		

PCTWORLD INTELLECTUAL PROPERTY ORGANIZATION
International Bureau

INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification ⁷ : C12N 15/13, 15/63, C07K 16/00, A61K 39/395, C12N 5/10		A1	(11) International Publication Number: WO 00/29584 (43) International Publication Date: 25 May 2000 (25.05.00)
(21) International Application Number: PCT/US99/27153 (22) International Filing Date: 16 November 1999 (16.11.99) (30) Priority Data: 60/108,945 18 November 1998 (18.11.98) US (71) Applicant: GENENTECH, INC. [US/US]; 1 DNA Way, South San Francisco, CA 94080-4990 (US). (72) Inventors: CHEN, Yvonne, M.; 1951 O'Farrell Street #321, San Mateo, CA 94403 (US). LOWMAN, Henry, B.; 400 San Juan Avenue, P.O. Box 2556, El Granada, CA 94018 (US). MULLER, Yves; Steenerbusch-Strasse 74, D-16341 Zepernick (DE). (74) Agents: WENDY, M., Lee et al.; Genentech, Inc., 1 DNA Way, South San Francisco, CA 94080-4990 (US).			(81) Designated States: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG). Published <i>With international search report. Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.</i>
(54) Title: ANTIBODY VARIANTS WITH HIGHER BINDING AFFINITY COMPARED TO PARENT ANTIBODIES			
(57) Abstract Antibody variants of parent antibodies are disclosed which have one or more amino acids inserted in a hypervariable region of the parent antibody and a binding affinity for a target antigen which is at least about two fold stronger than the binding affinity of the parent antibody for the antigen.			

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
AU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav Republic of Macedonia	TM	Turkmenistan
BF	Burkina Faso	GR	Greece	ML	Mali	TR	Turkey
BG	Bulgaria	HU	Hungary	MN	Mongolia	TT	Trinidad and Tobago
BJ	Benin	IE	Ireland	MR	Mauritania	UA	Ukraine
BR	Brazil	IL	Israel	MW	Malawi	UG	Uganda
BY	Belarus	IS	Iceland	MX	Mexico	US	United States of America
CA	Canada	IT	Italy	NE	Niger	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NL	Netherlands	VN	Viet Nam
CG	Congo	KE	Kenya	NO	Norway	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NZ	New Zealand	ZW	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's Republic of Korea	PL	Poland		
CM	Cameroon	KR	Republic of Korea	PT	Portugal		
CN	China	KZ	Kazakhstan	RO	Romania		
CU	Cuba	LC	Saint Lucia	RU	Russian Federation		
CZ	Czech Republic	LI	Liechtenstein	SD	Sudan		
DE	Germany	LK	Sri Lanka	SE	Sweden		
DK	Denmark	LR	Liberia	SG	Singapore		
EE	Estonia						

Description

5

10

15

20

25

30

35

40

45

50

55

ANTIBODY VARIANTS WITH HIGHER BINDING AFFINITY COMPARED TO PARENT ANTIBODIES

BACKGROUND OF THE INVENTION

Field of the Invention

This invention relates generally to antibody variants. In particular, antibody variants of parent antibodies are disclosed which have one or more amino acids inserted in a hypervariable region of the parent antibody and a binding affinity for a target antigen which is at least about two fold stronger than the binding affinity of the parent antibody for the antigen.

Description of Related Art

Antibodies are proteins, which exhibit binding specificity to a specific antigen. Native antibodies are usually heterotetrameric glycoproteins of about 150,000 daltons, composed of two identical light (L) chains and two identical heavy (H) chains. Each light chain is linked to a heavy chain by one covalent disulfide bond, while the number of disulfide linkages varies between the heavy chains of different immunoglobulin isotypes. Each heavy and light chain also has regularly spaced intrachain disulfide bridges. Each heavy chain has at one end a variable domain (V_H) followed by a number of constant domains. Each light chain has a variable domain at one end (V_L) and a constant domain at its other end; the constant domain of the light chain is aligned with the first constant domain of the heavy chain, and the light chain variable domain is aligned with the variable domain of the heavy chain. Particular amino acid residues are believed to form an interface between the light and heavy chain variable domains.

The term "variable" refers to the fact that certain portions of the variable domains differ extensively in sequence among antibodies and are responsible for the binding specificity of each particular antibody for its particular antigen. However, the variability is not evenly distributed through the variable domains of antibodies. It is concentrated in three segments called Complementarity Determining Regions (CDRs) both in the light chain and the heavy chain variable domains. The more highly conserved portions of the variable domains are called the framework regions (FR). The variable domains of native heavy and light chains each comprise four FR regions, largely adopting a β -sheet configuration, connected by

three CDRs, which form loops connecting, and in some cases forming part of, the β -sheet structure. The CDRs in each chain are held together in close proximity by the FR regions and, with the CDRs from the other chain, contribute to the formation of the antigen binding site of antibodies (see Kabat et al., *Sequences of Proteins of Immunological Interest*, 5th Ed. Public Health Service, National Institutes of Health, Bethesda, MD. (1991)).

The constant domains are not involved directly in binding an antibody to an antigen, but exhibit various effector functions. Depending on the amino acid sequence of the constant region of their heavy chains, antibodies or immunoglobulins can be assigned to different classes. There are five major classes of immunoglobulins: IgA, IgD, IgE, IgG and IgM, and several of these may be further divided into subclasses (isotypes), e.g. IgG1, IgG2, IgG3, and IgG4; IgA1 and IgA2. The heavy chain constant regions that correspond to the different classes of immunoglobulins are called α , δ , ϵ , γ , and μ , respectively. Of the various human immunoglobulin classes, only human IgG1, IgG2, IgG3 and IgM are known to activate complement.

In vivo, affinity maturation of antibodies is driven by antigen selection of higher affinity antibody variants which are made primarily by somatic hypermutagenesis. A "repertoire shift" also often occurs in which the predominant germline genes of the secondary or tertiary response are seen to differ from those of the primary or secondary response.

Various research groups have attempted to mimic the affinity maturation process of the immune system, by introducing mutations into antibody genes in vitro and using affinity selection to isolate mutants with improved affinity. Such mutant antibodies can be displayed on the surface of filamentous bacteriophage and antibodies can be selected by their affinity for antigen or by their kinetics of dissociation (off-rate) from antigen. Hawkins et al. *J. Mol. Biol.* 226:889-896 (1992). CDR walking mutagenesis has been employed to affinity mature human antibodies which bind the human envelope glycoprotein gp120 of human immunodeficiency virus type 1 (HIV-1) (Barbas III et al. *PNAS (USA)* 91: 3809-3813 (1994); and Yang et al. *J. Mol. Biol.* 254:392-403 (1995)); and an anti-c-erbB-2 single chain Fv fragment (Schier et al. *J. Mol. Biol.* 263:551567 (1996)). Antibody chain shuffling and CDR mutagenesis were used to affinity mature a high-affinity human antibody directed against the third hypervariable loop of HIV (Thompson et al. *J. Mol. Biol.* 256:77-88 (1996)). Balint and

5 Larrick Gene 137:109-118 (1993) describe a technique they coin
"parsimonious mutagenesis" which involves computer-assisted
oligodeoxyribonucleotide-directed scanning mutagenesis whereby all three
10 CDRs of a variable region gene are simultaneously and thoroughly searched
5 for improved variants. Wu et al. affinity matured an $\alpha\text{v}\beta 3$ -specific
humanized antibody using an initial limited mutagenesis strategy in which
every position of all six CDRs was mutated followed by the expression and
15 screening of a combinatorial library including the highest affinity mutants
(Wu et al. *FNAS (USA)* 95: 6037-6-42 (1998)). Phage antibodies are reviewed
10 in Chiswell and McCafferty *TIBTECH* 10:80-84 (1992); and Rader and Barbas
III *Current Opinion in Biotech.* 8:503-508 (1997). In each case where
20 mutant antibodies with improved affinity compared to a parent antibody are
reported in the above references, the mutant antibody has amino acid
substitutions in a CDR.

15

SUMMARY OF THE INVENTION

25 Unlike the affinity matured antibodies of the above references, the
present invention provides an antibody variant of a parent antibody, which
antibody variant comprises an amino acid insertion in or adjacent to a
30 hypervariable region of the parent antibody and has a binding affinity for
20 a target antigen which is at least about two fold stronger than the binding
affinity of the parent antibody for the antigen.

35 The invention further provides an antibody variant comprising a heavy
chain variable domain, wherein CDR H3 of the heavy chain variable domain
25 comprises the amino acid sequence of CDR H3 of a variant selected from the
group consisting of Y0239-19 (SEQ ID NO:85); Y0239-8 (SEQ ID NO:53); Y0240-
1 (SEQ ID NO:86); Y0239-12 (SEQ ID NO:78); Y0239-9 (SEQ ID NO:54); and
40 Y0261-6 (SEQ ID NO:89). These CDR H3 sequences may, for example, be
provided in the heavy chain variable domain sequence of SEQ ID NO: 98 or
30 99; see Figure 1B). Preferably, the antibody variant further comprises a
light chain variable domain and binds VEGF antigen with stronger binding
affinity than Y0192 (see Figures 1A and 1B; SEQ ID NO's 95 and 96).

45 The invention further provides a method for producing an antibody
variant comprising introducing an amino acid residue in or adjacent to a
35 hypervariable region of a parent antibody, wherein the antibody variant has
a binding affinity for a target antigen which is at least about two fold
50 stronger than the binding affinity of the parent antibody for said antigen.

5
Additionally, the invention provides a method for making an antibody variant, comprising the steps of:

10 (a) identifying potential amino acid interactions between a hypervariable region of a parent antibody and a target antigen;

5 (b) preparing a variant of the parent antibody comprising introducing an amino acid residue in or adjacent to the hypervariable region of the parent antibody, wherein the introduced amino acid residue contributes to the potential amino acid interactions in (a); and

15 (c) selecting an antibody variant prepared as in (b) which has a stronger binding affinity for the antigen than the parent antibody.

20 Various forms of the antibody variant are contemplated herein. For example, the antibody variant may be a full length antibody (e.g. having a human immunoglobulin constant region) or an antibody fragment (e.g. a F(ab')₂). Furthermore, the antibody variant may be labeled with a detectable label, immobilized on a solid phase and/or conjugated with a heterologous compound (such as a cytotoxic agent).

25 Diagnostic and therapeutic uses for the antibody variant are contemplated. In one diagnostic application, the invention provides a method for determining the presence of an antigen of interest comprising exposing a sample suspected of containing the antigen to the antibody variant and determining binding of the antibody variant to the sample. For this use, the invention provides a kit comprising the antibody variant and instructions for using the antibody variant to detect the antigen.

30 The invention further provides: isolated nucleic acid encoding the antibody variant; a vector comprising the nucleic acid, optionally, operably linked to control sequences recognized by a host cell transformed with the vector; a host cell transformed with the vector; a process for producing the antibody variant comprising culturing this host cell so that the nucleic acid is expressed and, optionally, recovering the antibody variant from the host cell culture (e.g. from the host cell culture medium).

35 The invention also provides a composition comprising the antibody variant and a pharmaceutically acceptable carrier or diluent. This composition for therapeutic use is sterile and may be lyophilized.

40 The invention further provides a method for treating a mammal comprising administering an effective amount of the antibody variant to the mammal.

5

Brief Description of the Drawings

10

Figures 1A and 1B show a sequence alignment of the light chain variable region (Figure 1A) and heavy chain variable region (Figure 1B) of several variants of the humanized anti-VEGF antibody F(ab)-12. The parental Fab-phage clone Y0192 contains light chain mutations which do not significantly affect antigen binding affinity, and has been described (WO98/45331). Another variant, Y0238-3, contains mutations in CDR H1 which improve antigen-binding (WO98/45331). Variant Y0239-19 contains the "VNERK" motif identified in selections from CDR H3 insertion libraries described herein. Variant Y0313-2 contains the CDR H1 mutations of Y0238-3 combined with the CDR H3 mutations of Y0239-19. Differences from F(ab)-12 are highlighted with shaded boxes. The sequence identifiers in Figures 1A and 1B are as follows: F(ab)-12 light chain variable domain (SEQ ID NO:94); Y0192, Y0238-3, Y0239-19 and Y0313-2 light chain variable domain (SEQ ID NO:95); F(ab)-12 and Y0192 heavy chain variable domain (SEQ ID NO:96); Y0238-3 heavy chain variable domain (SEQ ID NO:97); Y0239-19 heavy chain variable domain (SEQ ID NO:98); and Y0313-2 heavy chain variable domain (SEQ ID NO:99).

20

25

30

Figure 2 shows the inhibition of VEGF activity in a cell-based bioassay by Fab, F(ab)-12 and Fab variant Y0313-2.

35

Figure 3 shows a portion of the three-dimensional model of F(ab)-12 in complex with VEGF as determined by x-ray crystallography (Muller et al. Structure 6(9): 1153-1167 (1998)). The main chain trace of the CDR H3 region of the antibody is depicted as a magenta ribbon at right. A surface rendering of a portion of VEGF is depicted at left, with several proximal residues highlighted in red (acidic) or purple (basic). The side chain of D41 of VEGF can be seen as a site of potential interaction with a hypothetical insertion peptide placed into the CDR H3.

40

Figure 4 shows a superposition of portions of the three-dimensional model of F(ab)-12 in complex with VEGF (both molecules in gray; Muller et al., supra) with a model of the insertion variant Fab Y0313-2 (green) in complex with VEGF (yellow). The latter model is based on x-ray crystallographic determination of the variant complex structure described herein. The figure illustrates that little structural change is observed in the complex as compared with the F(ab)-12 complex, except in the immediate vicinity of the mutations V104, N104a, E104b, R104c, and K105.

45

50

Figure 5 shows a comparison of portions of the three-dimensional

5 model of F(ab)-12 in complex with VEGF (at right; Muller et al., supra)
with a model of Fab YG313-2 in complex with VEGF (at left) as described
herein. In each case, VEGF is shown in yellow, and the respective Fab is
10 shown in green. In the Y0313-2 complex, it can be seen that V104 and R104c
5 make new contacts with VEGF.

Detailed Description of the Preferred Embodiments

I. Definitions

15 The term "antibody" is used in the broadest sense and specifically
10 covers monoclonal antibodies (including full length monoclonal antibodies),
polyclonal antibodies, multispecific antibodies (e.g., bispecific
20 antibodies), and antibody fragments so long as they exhibit the desired
biological activity.

25 The term "hypervariable region" when used herein refers to the
15 regions of an antibody variable domain which are hypervariable in sequence
and/or form structurally defined loops. The hypervariable region comprises
amino acid residues from a "complementarity determining region" or "CDR"
(i.e. residues 24-34 ("CDR L1"), 50-56 ("CDR L2") and 89-97 ("CDR L3") in
the light chain variable domain and 31-35 ("CDR H1"), 50-65 ("CDR H2") and
20 95-102 ("CDR H3") in the heavy chain variable domain; Kabat et al.,
Sequences of Proteins of Immunological Interest, 5th Ed. Public Health
Service, National Institutes of Health, Bethesda, MD. (1991)) and/or those
residues from a "hypervariable loop" (i.e. residues 26-32 ("loop L1"), 50-
52 ("loop L2") and 91-96 ("loop L3") in the light chain variable domain and
35 26-32 ("loop H1"), 53-55 ("loop H2") and 96-101 ("loop H3") in the heavy
chain variable domain; Chothia and Lesk J. Mol. Biol. 196:901-917 (1987)).
In both cases, the variable domain residues are numbered according to Kabat
et al., supra. "Framework" or "FR" residues are those variable domain
40 residues other than the hypervariable region residues as herein defined.

30 The expression "variable domain residue numbering as in Kabat" refers
to the numbering system used for heavy chain variable domains or light
chain variable domains of the compilation of antibodies in Kabat et al.,
45 Sequences of Proteins of Immunological Interest, 5th Ed. Public Health
Service, National Institutes of Health, Bethesda, MD. (1991). Using this
35 numbering system, the actual linear amino acid sequence may contain fewer
or additional amino acids corresponding to a shortening of, or insertion
50 into, a FR or CDR of the variable domain. For example, a heavy chain

5 variable domain may include a single amino acid insert (residue 52a according to Kabat) after residue 52 of CDR H2 and inserted residues (e.g. residues 82a, 82b, and 82c, etc according to Kabat) after heavy chain FR
10 residue 82. The Kabat numbering of residues may be determined for a given antibody by alignment at regions of homology of the sequence of the
5 antibody with a "standard" Kabat numbered sequence.

"Antibody fragments" comprise a portion of a full length antibody, generally the antigen binding or variable region thereof. Examples of
15 antibody fragments include Fab, Fab', F(ab')₂, and Fv fragments; diabodies; linear antibodies; single-chain antibody molecules; and multispecific
10 antibodies formed from antibody fragments.

The term "monoclonal antibody" as used herein refers to an antibody obtained from a population of substantially homogeneous antibodies, i.e., the individual antibodies comprising the population are identical except
20 for possible naturally occurring mutations that may be present in minor amounts. Monoclonal antibodies are highly specific, being directed against a single antigenic site. Furthermore, in contrast to conventional
25 (polyclonal) antibody preparations which typically include different antibodies directed against different determinants (epitopes), each monoclonal antibody is directed against a single determinant on the
30 antigen. The modifier "monoclonal" indicates the character of the antibody as being obtained from a substantially homogeneous population of antibodies, and is not to be construed as requiring production of the antibody by any particular method. For example, the monoclonal antibodies
35 to be used in accordance with the present invention may be made by the hybridoma method first described by Kohler et al., *Nature* 256:495 (1975), or may be made by recombinant DNA methods (see, e.g., U.S. Patent No. 4,816,567). The "monoclonal antibodies" may also be isolated from phage
40 antibody libraries using the techniques described in Clackson et al., *Nature* 352:624-628 (1991) and Marks et al., *J. Mol. Biol.* 222:581-597 (1991), for example.

The monoclonal antibodies herein specifically include "chimeric" antibodies (immunoglobulins) in which a portion of the heavy and/or light
45 chain is identical with or homologous to corresponding sequences in antibodies derived from a particular species or belonging to a particular
35 antibody class or subclass, while the remainder of the chain(s) is identical with or homologous to corresponding sequences in antibodies
50

5 derived from another species or belonging to another antibody class or subclass, as well as fragments of such antibodies, so long as they exhibit the desired biological activity (U.S. Patent No. 4,816,567; and Morrison et al., *Proc. Natl. Acad. Sci. USA* 81:6851-6855 (1984)).

10 "Humanized" forms of non-human (e.g., murine) antibodies are chimeric antibodies which contain minimal sequence derived from non-human immunoglobulin. For the most part, humanized antibodies are human immunoglobulins (recipient antibody) in which residues from a hypervariable region of the recipient are replaced by residues from a hypervariable region of a non-human species (donor antibody) such as mouse, rat, rabbit or nonhuman primate having the desired specificity, affinity, and capacity. In some instances, Fv framework region (FR) residues of the human immunoglobulin are replaced by corresponding non-human residues. Furthermore, humanized antibodies may comprise residues which are not found in the recipient antibody or in the donor antibody. These modifications are made to further refine antibody performance. In general, the humanized antibody will comprise substantially all of at least one, and typically two, variable domains, in which all or substantially all of the hypervariable loops correspond to those of a non-human immunoglobulin and all or substantially all of the FR regions are those of a human immunoglobulin sequence. The humanized antibody optionally also will comprise at least a portion of an immunoglobulin constant region (Fc), typically that of a human immunoglobulin. For further details, see Jones et al., *Nature* 321:522-525 (1986); Riechmann et al., *Nature* 332:323-329 (1988); and Presta, *Curr. Op. Struct. Biol.* 2:593-596 (1992).

35 "Single-chain Fv" or "sFv" antibody fragments comprise the V_H and V_L domains of antibody, wherein these domains are present in a single polypeptide chain. Generally, the Fv polypeptide further comprises a polypeptide linker between the V_H and V_L domains which enables the sFv to form the desired structure for antigen binding. For a review of sFv see Pluckthun in *The Pharmacology of Monoclonal Antibodies*, vol. 113, Rosenberg and Moore eds. Springer-Verlag, New York, pp. 269-315 (1994).

45 The term "diabodies" refers to small antibody fragments with two antigen-binding sites, which fragments comprise a heavy chain variable domain (V_H) connected to a light chain variable domain (V_L) in the same polypeptide chain (V_H - V_L). By using a linker that is too short to allow pairing between the two domains on the same chain, the domains are forced

5 to pair with the complementary domains of another chain and create two antigen-binding sites. Diabodies are described more fully in, for example, EP 404,097; WO 93/11161; and Hollinger et al., *Proc. Natl. Acad. Sci. USA* 90:6444-6448 (1993).

10 The expression "linear antibodies" when used throughout this application refers to the antibodies described in Zapata et al. *Protein Eng.* 8(10):1057-1062 (1995). Briefly, these antibodies comprise a pair of tandem Fd segments ($V_H\text{-}C_H1\text{-}V_H\text{-}C_H1$) which form a pair of antigen binding regions. Linear antibodies can be bispecific or monospecific.

15 A "parent antibody" is an antibody comprising an amino acid sequence which lacks, or is deficient in, one or more amino acid residues in or adjacent to one or more hypervariable regions thereof compared to an antibody variant as herein disclosed. Thus, the parent antibody has a shorter hypervariable region than the corresponding hypervariable region of an antibody variant as herein disclosed. The parent polypeptide may comprise a native sequence (i.e. a naturally occurring) antibody (including a naturally-occurring allelic variant) or an antibody with pre-existing amino acid sequence modifications (such as other insertions, deletions and/or substitutions) of a naturally-occurring sequence. Preferably the parent antibody is a humanized antibody or a human antibody.

20 As used herein, "antibody variant" refers to an antibody which has an amino acid sequence which differs from the amino acid sequence of a parent antibody. Preferably, the antibody variant comprises a heavy chain variable domain or a light chain variable domain having an amino acid sequence which is not found in nature. Such variants necessarily have less than 100% sequence identity or similarity with the parent antibody. In a preferred embodiment, the antibody variant will have an amino acid sequence from about 75% to less than 100% amino acid sequence identity or similarity with the amino acid sequence of either the heavy or light chain variable domain of the parent antibody, more preferably from about 80% to less than 100%, more preferably from about 85% to less than 100%, more preferably from about 90% to less than 100%, and most preferably from about 95% to less than 100%. Identity or similarity with respect to this sequence is defined herein as the percentage of amino acid residues in the candidate sequence that are identical (i.e. same residue) with the parent antibody residues, after aligning the sequences and introducing gaps, if necessary, to achieve the maximum percent sequence identity. None of N-terminal, C-

terminal, or internal extensions, deletions, or insertions into the antibody sequence outside of the variable domain shall be construed as affecting sequence identity or similarity. The antibody variant is generally one which has a longer hypervariable region (by one or more amino acid residues; e.g. by about one to about 30 amino acid residues and preferably by about two to about ten amino acid residues) than the corresponding hypervariable region of a parent antibody.

An "amino acid alteration" refers to a change in the amino acid sequence of a predetermined amino acid sequence. Exemplary alterations include insertions, substitutions and deletions.

An "amino acid insertion" refers to the introduction of one or more amino acid residues into a predetermined amino acid sequence.

The amino acid insertion may comprise a "peptide insertion" in which case a peptide comprising two or more amino acid residues joined by peptide bond(s) is introduced into the predetermined amino acid sequence. Where the amino acid insertion involves insertion of a peptide, the inserted peptide may be generated by random mutagenesis such that it has an amino acid sequence which does not exist in nature.

The inserted residue or residues may be "naturally occurring amino acid residues" (i.e. encoded by the genetic code) and selected from the group consisting of: alanine (Ala); arginine (Arg); asparagine (Asn); aspartic acid (Asp); cysteine (Cys); glutamine (Gln); glutamic acid (Glu); glycine (Gly); histidine (His); isoleucine (Ile); leucine (Leu); lysine (Lys); methionine (Met); phenylalanine (Phe); proline (Pro); serine (Ser); threonine (Thr); tryptophan (Trp); tyrosine (Tyr); and valine (Val).

Insertion of one or more non-naturally occurring amino acid residues is also encompassed by the definition of an amino acid insertion herein. A "non-naturally occurring amino acid residue" refers to a residue, other than those naturally occurring amino acid residues listed above, which is able to covalently bind adjacent amino acid residues(s) in a polypeptide chain. Examples of non-naturally occurring amino acid residues include norleucine, ornithine, norvaline, homoserine and other amino acid residue analogues such as those described in Ellman et al. *Meth. Enzym.* 202:301-336 (1991). To generate such non-naturally occurring amino acid residues, the procedures of Noren et al. *Science* 244:182 (1989) and Ellman et al., *supra*, can be used. Briefly, these procedures involve chemically activating a suppressor tRNA with a non-naturally occurring amino acid residue followed

5 by *in vitro* transcription and translation of the RNA.

10 An amino acid insertion "in a hypervariable region" refers to the introduction of one or more amino acid residues within a hypervariable region amino acid sequence.

15 An amino acid insertion "adjacent a hypervariable region" refers to the introduction of one or more amino acid residues at the N-terminal and/or C-terminal end of a hypervariable region, such that at least one of the inserted amino acid residues forms a peptide bond with the N-terminal or C-terminal amino acid residue of the hypervariable region in question.

20 An "amino acid substitution" refers to the replacement of an existing amino acid residue in a predetermined amino acid sequence with another different amino acid residue.

25 The term "potential amino acid interactions" refers to contacts or energetically favorable interactions between one or more amino acid residues present in an antigen and one or more amino acid residues which do not exist in a parent antibody but can be introduced therein so as to increase the amino acid contacts between the antigen and an antibody variant comprising those introduced amino acid residue(s). Preferably the amino acid interactions of interest are selected from the group consisting of hydrogen bonding, van der Waals interactions and ionic interactions.

30 The term "target antigen" herein refers to a predetermined antigen to which both a parent antibody and antibody variant as herein defined bind. The target antigen may be polypeptide, carbohydrate, nucleic acid, lipid, hapten or other naturally occurring or synthetic compound. Preferably, the target antigen is a polypeptide. While the antibody variant binds the target antigen with better binding affinity than the parent antibody, the parent antibody generally has a binding affinity (K_d) value for the target antigen of no more than about $1 \times 10^{-5} M$, and preferably no more than about $1 \times 10^{-6} M$.

35 An "isolated" antibody is one which has been identified and separated and/or recovered from a component of its natural environment. Contaminant components of its natural environment are materials which would interfere with diagnostic or therapeutic uses for the antibody, and may include enzymes, hormones, and other proteinaceous or nonproteinaceous solutes. In preferred embodiments, the antibody will be purified (1) to greater than 95% by weight of antibody as determined by the Lowry method, and most

preferably more than 99% by weight, (2) to a degree sufficient to obtain at least 15 residues of N-terminal or internal amino acid sequence by use of a spinning cup sequenator, or (3) to homogeneity by SDS-PAGE under reducing or nonreducing conditions using Coomassie blue or, preferably, silver stain. Isolated antibody includes the antibody in situ within recombinant cells since at least one component of the antibody's natural environment will not be present. Ordinarily, however, isolated antibody will be prepared by at least one purification step.

"Treatment" refers to both therapeutic treatment and prophylactic or preventative measures. Those in need of treatment include those already with the disorder as well as those in which the disorder is to be prevented.

A "disorder" is any condition that would benefit from treatment with the antibody variant. This includes chronic and acute disorders or diseases including those pathological conditions which predispose the mammal to the disorder in question.

"Mammal" for purposes of treatment refers to any animal classified as a mammal, including humans, domestic and farm animals, nonhuman primates, and zoo, sports, or pet animals, such as dogs, horses, cats, cows, etc.

The term "cytotoxic agent" as used herein refers to a substance that inhibits or prevents the function of cells and/or causes destruction of cells. The term is intended to include radioactive isotopes (e.g., I^{131} , I^{125} , Y^{90} and Re^{186}), chemotherapeutic agents, and toxins such as enzymatically active toxins of bacterial, fungal, plant or animal origin, or fragments thereof.

A "chemotherapeutic agent" is a chemical compound useful in the treatment of cancer. Examples of chemotherapeutic agents include Adriamycin, Doxorubicin, 5-Fluorouracil, Cytosine arabinoside ("Ara-C"), Cyclophosphamide, Thiotepa, Taxotere (docetaxel), Busulfan, Cytosin, Taxol, Methotrexate, Cisplatin, Melphalan, Vinblastine, Bleomycin, Etoposide, Ifosfamide, Mitomycin C, Mitoxantrone, Vincristine, Vinorelbine, Carboplatin, Teniposide, Daunomycin, Carminomycin, Aminopterin, Dactinomycin, Mitomycins, Esperamicins (see U.S. Pat. No. 4,675,137), Melphalan and other related nitrogen mustards.

The term "prodrug" as used in this application refers to a precursor or derivative form of a pharmaceutically active substance that is less cytotoxic to tumor cells compared to the parent drug and is capable of

5 being enzymatically activated or converted into the more active parent form. See, e.g., Wilman, "Prodrugs in Cancer Chemotherapy" *Biochemical Society Transactions*, 14, pp. 375-382, 615th Meeting Belfast (1986) and
10 Stella et al., "Prodrugs: A Chemical Approach to Targeted Drug Delivery,"
5 *Directed Drug Delivery*, Borchardt et al., (ed.), pp. 247-267, Humana Press (1985). The prodrugs of this invention include, but are not limited to, phosphate-containing prodrugs, thiophosphate-containing prodrugs, sulfate-containing prodrugs, peptide-containing prodrugs, D-amino acid-modified prodrugs, glycosylated prodrugs, beta-lactam-containing prodrugs,
15 optionally substituted phenoxyacetamide-containing prodrugs or optionally substituted phenylacetamide-containing prodrugs, 5-fluorocytosine and other 5-fluorouridine prodrugs which can be converted into the more active cytotoxic free drug. Examples of cytotoxic drugs that can be derivatized into a prodrug form for use in this invention include, but are not limited
20 to, those chemotherapeutic agents described above.

25 The word "label" when used herein refers to a detectable compound or composition which is conjugated directly or indirectly to the antibody. The label may be itself be detectable (e.g., radioisotope labels or fluorescent labels) or, in the case of an enzymatic label, may catalyze
30 chemical alteration of a substrate compound or composition which is detectable.

By "solid phase" is meant a non-aqueous matrix to which the antibody of the present invention can adhere. Examples of solid phases encompassed herein include those formed partially or entirely of glass (e.g. controlled
35 pore glass), polysaccharides (e.g., agarose), polyacrylamides, polystyrene, polyvinyl alcohol and silicones. In certain embodiments, depending on the context, the solid phase can comprise the well of an assay plate; in others it is a purification column (e.g. an affinity chromatography column). This
40 term also includes a discontinuous solid phase of discrete particles, such as those described in U.S. Patent No. 4,275,149.

45 A "liposome" is a small vesicle composed of various types of lipids, phospholipids and/or surfactant which is useful for delivery of a drug (such as the antibody variants disclosed herein and, optionally, a chemotherapeutic agent) to a mammal. The components of the liposome are
35 commonly arranged in a bilayer formation, similar to the lipid arrangement of biological membranes.

50 An "isolated" nucleic acid molecule is a nucleic acid molecule that

5 is identified and separated from at least one contaminant nucleic acid molecule with which it is ordinarily associated in the natural source of the antibody nucleic acid. An isolated nucleic acid molecule is other than
10 in the form or setting in which it is found in nature. Isolated nucleic acid molecules therefore are distinguished from the nucleic acid molecule as it exists in natural cells. However, an isolated nucleic acid molecule includes a nucleic acid molecule contained in cells that ordinarily express the antibody where, for example, the nucleic acid molecule is in a
15 chromosomal location different from that of natural cells.

10 The expression "control sequences" refers to DNA sequences necessary for the expression of an operably linked coding sequence in a particular host organism. The control sequences that are suitable for prokaryotes, for example, include a promoter, optionally an operator sequence, and a ribosome binding site. Eukaryotic cells are known to utilize promoters,
20 polyadenylation signals, and enhancers.

25 Nucleic acid is "operably linked" when it is placed into a functional relationship with another nucleic acid sequence. For example, DNA for a presequence or secretory leader is operably linked to DNA for a polypeptide if it is expressed as a preprotein that participates in the secretion of the polypeptide; a promoter or enhancer is operably linked to a coding
30 sequence if it affects the transcription of the sequence; or a ribosome binding site is operably linked to a coding sequence if it is positioned so as to facilitate translation. Generally, "operably linked" means that the DNA sequences being linked are contiguous, and, in the case of a secretory
35 leader, contiguous and in reading phase. However, enhancers do not have to be contiguous. Linking is accomplished by ligation at convenient restriction sites. If such sites do not exist, the synthetic oligonucleotide adaptors or linkers are used in accordance with
40 conventional practice.

30 As used herein, the expressions "cell," "cell line," and "cell culture" are used interchangeably and all such designations include progeny. Thus, the words "transformants" and "transformed cells" include
45 the primary subject cell and cultures derived therefrom without regard for the number of transfers. It is also understood that all progeny may not be precisely identical in DNA content, due to deliberate or inadvertent
35 mutations. Mutant progeny that have the same function or biological activity as screened for in the originally transformed cell are included.
50

Where distinct designations are intended, it will be clear from the context.

II. Modes for Carrying out the Invention

The invention herein relates to a method for making an antibody variant. The parent antibody or starting antibody is prepared using techniques available in the art for generating such antibodies. Exemplary methods for generating antibodies are described in more detail in the following sections.

The parent antibody is directed against a target antigen of interest. Preferably, the target antigen is a biologically important polypeptide and administration of the antibody to a mammal suffering from a disease or disorder can result in a therapeutic benefit in that mammal. However, antibodies directed against nonpolypeptide antigens (such as tumor-associated glycolipid antigens; see US Patent 5,091,178) are also contemplated.

Where the antigen is a polypeptide, it may be a transmembrane molecule (e.g. receptor) or ligand such as a growth factor. Exemplary antigens include molecules such as renin; a growth hormone, including human growth hormone and bovine growth hormone; growth hormone releasing factor; parathyroid hormone; thyroid stimulating hormone; lipoproteins; alpha-1-antitrypsin; insulin A-chain; insulin B-chain; proinsulin; follicle stimulating hormone; calcitonin; luteinizing hormone; glucagon; clotting factors such as factor VIIIC, factor IX, tissue factor, and von Willebrands factor; anti-clotting factors such as Protein C; atrial natriuretic factor; lung surfactant; a plasminogen activator, such as urokinase or human urine or tissue-type plasminogen activator (t-PA); bombesin; thrombin; hemopoietic growth factor; tumor necrosis factor-alpha and -beta; enkephalinase; RANTES (regulated on activation normally T-cell expressed and secreted); human macrophage inflammatory protein (MIP-1-alpha); a serum albumin such as human serum albumin; Muellerian-inhibiting substance; relaxin A-chain; relaxin B-chain; prorelaxin; mouse gonadotropin-associated peptide; a microbial protein, such as beta-lactamase; DNase; IgE; a cytotoxic T-lymphocyte associated antigen (CTLA), such as CTLA-4; inhibin; activin; vascular endothelial growth factor (VEGF); receptors for hormones or growth factors; protein A or D; rheumatoid factors; a neurotrophic factor such as bone-derived neurotrophic factor (BDNF), neurotrophin-3, -4, -5, or -6 (NT-3, NT-4, NT-5, or NT-6), or a nerve growth factor; platelet-

5 derived growth factor (PDGF); fibroblast growth factor such as aFGF and bFGF; epidermal growth factor (EGF); transforming growth factor (TGF) such as TGF-alpha and TGF-beta; insulin-like growth factor-I and -II (IGF-I and IGF-II); des(1-3)-IGF-I (brain IGF-I), insulin-like growth factor binding
10 proteins; CD proteins such as CD3, CD4, CD8, CD19 and CD20; erythropoietin; osteoinductive factors; immunotoxins; a bone morphogenetic protein (BMP); an interferon such as interferon-alpha, -beta, and -gamma; colony
15 stimulating factors (CSFs), e.g., M-CSF, GM-CSF, and G-CSF; interleukins (ILs), e.g., IL-1 to IL-10; superoxide dismutase; T-cell receptors; surface
20 membrane proteins; decay accelerating factor; viral antigen such as, for example, a portion of the AIDS envelope; transport proteins; homing receptors; addressins; regulatory proteins; integrins such as CD11a, CD11b, CD11c, CD18, an ICAM, VLA-4 and VCAM; a tumor associated antigen such as HER2, HER3 or HER4 receptor; and fragments of any of the above-listed
25 polypeptides.

Preferred molecular targets for antibodies encompassed by the present invention include CD proteins such as CD3, CD4, CD8, CD19, CD20 and CD34; members of the ErbB receptor family such as the EGF receptor, HER2, HER3 or HER4 receptor; cell adhesion molecules such as LFA-1, Mac1, p150,95, VLA-4, ICAM-1, VCAM and $\alpha v/\beta 3$ integrin including either alpha or
30 beta subunits thereof (e.g. anti-CD11a, anti-CD18 or anti-CD11b antibodies); growth factors such as VEGF; IgE; blood group antigens; flk2/flt3 receptor; obesity (OB) receptor; mpl receptor; CTLA-4; protein C etc.

35 25 The antigen used to generate an antibody may be isolated from a natural source thereof, or may be produced recombinantly or made using other synthetic methods. Alternatively, cells comprising native or recombinant antigen can be used as immunogens for making antibodies.

40 The parent antibody may have pre-existing strong binding affinity for the target antigen. For example, the parent antibody may bind the antigen of interest with a binding affinity (K_d) value of no more than about 1×10^{-7} M, preferably no more than about 1×10^{-8} M and most preferably no more
45 than about 1×10^{-9} M.

Antibody "binding affinity" may be determined by equilibrium methods
35 (e.g. enzyme-linked immunoabsorbent assay (ELISA) or radioimmunoassay (RIA)), or kinetics (e.g. BIACORE™ analysis; see Example 1 below), for example.

5

Also, the antibody may be subjected to other "biological activity assays", e.g., in order to evaluate its "potency" or pharmacological activity and potential efficacy as a therapeutic agent. Such assays are known in the art and depend on the target antigen and intended use for the antibody. Examples include the keratinocyte monolayer adhesion assay and the mixed lymphocyte response (MLR) assay for CD11a (see WO98/23761); tumor cell growth inhibition assays (as described in WO 89/06692, for example); antibody-dependent cellular cytotoxicity (ADCC) and complement-mediated cytotoxicity (CDC) assays (US Patent 5,500,362); agonistic activity or hematopoiesis assays (see WO 95/27062); tritiated thymidine incorporation assay; and alamar blue assay to measure metabolic activity of cells in response to a molecule such as VEGF (See Example 1 below).

10

5

15

20

The amino acid sequence of the parent antibody is altered so as to generate an antibody variant which has a stronger binding affinity for the target antigen than the parent antibody. The antibody variant preferably has a binding affinity for the target antigen which is at least about two fold stronger (e.g. from about two fold to about 1000 fold or even to about 10,000 fold improved binding affinity), preferably at least about five fold stronger, and preferably at least about ten fold or 100 fold stronger, than the binding affinity of the parent antibody for the antigen. The enhancement in binding affinity desired or required may depend on the initial binding affinity of the parent antibody.

15

25

30

Where the assay used is a biological activity assay, the antibody variant preferably has a potency in the biological activity assay of choice which is at least about two fold greater (e.g. from about two fold to about 1000 fold or even to about 10,000 fold improved potency), preferably at least about 20 fold greater, more preferably at least about 50 fold greater, and sometimes at least about 100 fold or 200 fold greater, than the biological activity of the parent antibody in that assay.

35

25

40

To generate the antibody variant, one or more amino acid residues are introduced or inserted in or adjacent to one or more of the hypervariable regions of the parent antibody. Generally, one will insert one or more amino acid residues in a CDR of the parent antibody. The number of residues to be inserted may be from about one residue to about 30 amino acid residues, e.g. from about two to about ten amino acid residues. In deciding the number of residues to be inserted, one may take into account the range of lengths of the hypervariable region in question in known

30

45

35

50

55

5 antibodies. For example, for the first hypervariable region of a light
chain variable domain, the hypervariable region is preferably "CDR L1"
according to Kabat et al., supra, e.g. having an overall length from about
10 nine amino acid residues to about 20 residues, including the inserted amino
5 acid residue(s). With respect to the second hypervariable region of a light
chain variable domain, the hypervariable region is preferably "CDR L2"
according to Kabat et al., supra, e.g. having an overall length from about
15 five amino acid residues to about ten residues, including the inserted
amino acid residue(s). In relation to the third hypervariable region of a
10 light chain variable domain, the hypervariable region is preferably "CDR
L3" according to Kabat et al., supra, e.g. having an overall length from
about seven amino acid residues to about 20 residues, including the
20 inserted amino acid residue(s).

Preferably, the antibody variant has one or more amino acid residues
15 inserted in a hypervariable region of the heavy chain variable region, most
preferably CDR H3. If this hypervariable region is chosen, preferably the
25 inserted amino acid residues are between residue numbers 97 and 102 (e.g.,
adjacent to, and preferably C-terminal in sequence to, residue number 100)
of the heavy chain variable domain of the parent antibody, utilizing the
20 variable domain residue numbering as in Kabat.

In deciding upon the number of amino acid residues to insert, one may
take into account the desired length of the altered hypervariable region.
For example, for the first hypervariable region of a heavy chain variable
domain, the hypervariable region is preferably the stretch of residues from
35 the "loop H1" of Chothia et al, supra, combined with the stretch of
25 residues considered to constitute "CDR H1" according to Kabat et al.,
supra. Thus, this first hypervariable loop of the heavy chain variable
domain may have an overall length from about eight amino acid residues to
40 about 20 residues including the inserted amino acid residue(s). In relation
30 to the second hypervariable region of a heavy chain variable domain, the
hypervariable region is preferably "CDR H2" according to Kabat et al.,
supra, e.g. having an overall length from about 14 amino acid residues to
45 about 25 residues, including the inserted amino acid residue(s). Finally,
in relation to the third hypervariable region of a heavy chain variable
35 domain, the hypervariable region is preferably "CDR H3" according to Kabat
et al., supra, e.g. having an overall length from about six amino acid
50 residues to about 30 residues, including the inserted amino acid

5 residue(s).

Antibody variants with inserted amino acid residue(s) in a hypervariable region thereof may be prepared randomly, especially where the starting binding affinity of the parent antibody for the target antigen is such that randomly produced antibody variants can be readily screened. For example, phage display provides a convenient method of screening such random variants.

The invention also provides a more systematic method for making antibody variants. This method involves the following general steps, usually performed sequentially:

(a) identifying potential amino acid interactions between a hypervariable region of a parent antibody and a target antigen;

(b) preparing a variant of the parent antibody by introducing an amino acid residue in or adjacent to the hypervariable region of the parent antibody, wherein the introduced amino acid residue contributes to the potential amino acid interactions in (a); and

(c) selecting an antibody variant prepared as in (b) which has a stronger binding affinity for the antigen than the parent antibody.

According to step (a) of this method, one may analyze a molecular model of the parent antibody complexed with antigen. The molecular model may be obtained from an X-ray crystal or nuclear magnetic resonance (NMR) structure of this complex. See, e.g., Amit et al. *Science* 233:747-753 (1986); and Muller et al. *Structure* 6(9): 1153-1167 (1998)). Alternatively, computer programs can be used to create molecular models of antibody/antigen complexes (see, e.g., Levy et al. *Biochemistry* 28:7168-7175 (1989); Brucoleri et al. *Nature* 335: 564-568 (1998); and Chothia et al. *Science* 233: 755-758 (1986)), where a crystal structure is not available.

In the preferred method, one analyzes the molecular model of the antigen/antibody complex and identifies potential areas for increasing energetically favorable interactions between the antigen and a hypervariable region of the antibody. For example, one may identify potential polar interactions (e.g. ion pairs and/or hydrogen-bonding); non-polar interactions (such as Van der Waals attractions and/or hydrophobic interactions); and/or covalent interactions (e.g. disulfide bond(s)) between one or more amino acid residues of the antigen and one or more amino acid residues which can be inserted in or adjacent to a hypervariable

5 region of the antibody. Preferably at least one of the inserted residues
has a net positive charge or a net negative charge. For example, at least
one of the inserted residues may be a positively charged residue,
10 preferably arginine or lysine.

5 Examples of side chains typically having positive charge are lysine,
arginine, and histidine. Examples of side chains typically having negative
charge are aspartic acid and glutamic acid. These side chains may undergo
15 ionic interactions (positive residues paired with negative residues), as
well as polar interactions with side chains having polar functional groups:
10 tryptophan, serine, threonine, tyrosine, cysteine, tyrosine, asparagine,
and glutamine. In addition, polar or ionic interactions may be mediated
through intervening solvent (such as water) or solute (e.g. phosphate or
20 sulfate) molecules.

Examples of residues which may be involved in hydrophobic
15 interactions, or non-polar Van der Waals interactions, are typically
alanine, valine, leucine, isoleucine, proline, phenylalanine, tryptophan,
25 methionine, and tyrosine; however, the non-polar side chains of other
residues, such as lysine or arginine, may also participate in such
interactions. Aromatic side chains such as phenylalanine, tyrosine, and
20 tryptophan may form aromatic (π) stacking interactions, or may act as
hydrogen-bond acceptors.

In addition, the main chain atoms of any residue (including glycine)
may undergo Van der Waals or hydrophobic interactions; and the atoms
nitrogen and carbonyl oxygen of the main chain, may undergo polar
35 (hydrogen-bonding) interactions. In some cases, a covalent bond
25 (disulfide) may be formed from a cysteine residue of the antibody with a
cysteine residue of the antigen.

Finally, post-translational modifications (e.g., glycosylation or
40 phosphorylation) or a prosthetic group (e.g., heme or zinc finger) may
30 provide additional functional groups (carboxylate or phosphate oxygens;
zinc or iron atoms) for interaction between antibody and antigen.

Thus, one may, for example, introduce one or more charged amino acid
45 residues in or adjacent to a hypervariable region of the parent antibody in
an appropriate three dimensional location, such that the introduced residue
35 or residues are able to form ion pair(s) with one or more oppositely
charged residues in the antigen. Similarly, one can create hydrogen-
50 bonding pair(s), Van der Waals interactions, etc., by introducing

appropriate amino acid residues in an appropriate location in or adjacent to a hypervariable region of the antibody.

The antibody variant may comprise additional alterations, such as amino acid deletions or substitutions in the hypervariable region of the antibody in which the insertion is made. This is shown in the example below, wherein the hypervariable region was modified by both amino acid substitutions as well as amino acid insertions.

In general, any inserted amino acid residue or inserted peptide will need to exit the existing antibody polypeptide chain at a residue position (x), extend to a point sufficiently near to the site of a new contact such that some portion of the amino acid side chain or main chain of the peptide can form an interaction, and return to reenter the existing antibody polypeptide chain at a position (y) (where $y > x$ in the linear sequence).

It is desirable that the inserted amino acid residue or peptide not significantly perturb the structure of the antibody in a global or local sense, beyond the vicinity of the newly inserted amino acid residue or peptide. In particular, the inserted amino acid residue or peptide preferably does not distort the FR residues of the antibody, or residues of the antibody or antigen involved in existing contacts. This may be evaluated in an actual or modeled complex.

If both exit/reentry residues (x and y) lack significant intramolecular and intermolecular contacts (i.e., both within the antibody, and between antibody and antigen), then an amino acid or peptide insertion may be accomplished by adding a peptide segment between residues x and y, leaving residues x and y unchanged. Alternatively, either or both residues x and y may be deleted and replaced by a peptide segment of >2 residues.

Often, residues x and y, and/or intervening residues in the parent antibody, may be involved in significant intramolecular and intermolecular contacts. In this case, these interactions may be maintained or replaced with residues contributing similar interactions, while allowing for an inserted residue or peptide to exit and reenter the chain. This may be accomplished by substituting the two residues x and y and/or intervening residues in the parent antibody with random residues, which can be subsequently subjected to affinity screening (or screening for other biological activities) to identify variants with improved affinity.

This systematic method is illustrated in Figure 3 for example, where residues D41 and E42 in the VEGF antigen were identified as potential

5 candidates for interacting with introduced residues in CDR H3 of the heavy chain variable domain of the parent antibody.

10 Thus, as illustrated in Figures 4 and 5, D41 of the VEGF antigen is able to form an ion pair with inserted residue R104c in CDR H3 of variant antibody Y0313-2 of the Example below. Figure 5 further shows how residue V104 in variant antibody Y0313-2 is able to form a hydrophobic interaction with residues 93 to 95 of the VEGF antigen. Thus, it can be seen that one identifies potential areas where the contacts between antigen and antibody can be improved, so as to increase the affinity of the antibody variant.

15 Generally one makes changes in hypervariable regions proximal to antigen when the antigen and antibody are complexed together. For example, the hypervariable region of the parent antibody which may be modified as disclosed herein generally has one or more amino acid residues within about 20 Å of one or more amino acid residues of the antigen. The hypervariable region to be altered herein may be one which, in the parent antibody, does not make significant contact with antigen (i.e. a non-contacting hypervariable region can be modified to become a contacting hypervariable region). Preferably however, the hypervariable region to be modified does contact antigen and the method herein serves to increase the contacts between the antigen and the already-contacting hypervariable region.

20 In another embodiment, one may identify hypervariable region residues which interact with antigen by alanine-scanning mutagenesis of the antigen and/or parent antibody (Muller et al. *Structure* 6(9): 1153-1167 (1998)) or by other means. Hypervariable regions identified as contacting antigen are candidates for amino acid insertion(s) as herein disclosed.

25 Nucleic acid molecules encoding amino acid sequence variants are prepared by a variety of methods known in the art. These methods include, but are not limited to, oligonucleotide-mediated (or site-directed) mutagenesis, PCR mutagenesis, and cassette mutagenesis of an earlier prepared variant or a non-variant version of the parent antibody. The preferred method for making variants is site directed mutagenesis (see, e.g., Kunkel, *Proc. Natl. Acad. Sci. USA* 82:488 (1985)). Moreover, a nucleic acid sequence can be made synthetically, once the desired amino acid sequence is arrived at conceptually. One can also make the antibody variant by peptide synthesis, peptide ligation or other methods.

30 Following production of the antibody variant, the activity of that molecule relative to the parent antibody may be determined. As noted

5 above, this may involve determining the binding affinity and/or other biological activities of the antibody. In a preferred embodiment of the invention, a panel of antibody variants are prepared and are screened for
10 binding affinity for the antigen and/or potency in one or more biological activity assays. One or more of the antibody variants selected from an initial screen are optionally subjected to one or more further biological
15 activity assays to confirm that the antibody variant(s) have improved activity in more than one assay.

One preferred method of making and screening insertion mutants
10 involves displaying antibody variants on the surface of filamentous bacteriophage and selecting antibody variants based on their affinity for antigen, by their kinetics of dissociation (off-rate) from antigen, or some
20 other screen for antibody affinity or potency. This was the method used to identify antibody variants with enhanced biological activity in the Example
15 below.

25 Aside from the above insertions in the hypervariable region of the parent antibody one may make other alterations in the amino acid sequences of one or more of the hypervariable regions. For example, the above amino acid insertions may be combined with deletions or substitutions of other
30 hypervariable region residues. Moreover, one or more alterations (e.g. substitutions) of FR residues may be introduced in the parent antibody where these result in an improvement in the binding affinity of the antibody variant for the antigen. Examples of framework region residues to
35 modify include those which non-covalently bind antigen directly (Amit et al. Science 233:747-753 (1986)); interact with/affect the conformation of a CDR (Chothia et al. J. Mol. Biol. 196:901-917 (1987)); and/or participate in the $V_L - V_H$ interface (EP 239 400B1). Such amino acid sequence alterations may be present in the parent antibody, may be made
40 simultaneously with the amino acid insertion(s) herein or may be made after a variant with an amino acid insertion is generated.
30

The antibody variants may be subjected to other modifications, oftentimes depending on the intended use of the antibody. Such
45 modifications may involve further alteration of the amino acid sequence, fusion to heterologous polypeptide(s) and/or covalent modification. With respect to amino acid sequence alterations, exemplary modifications are
35 elaborated above. For example, any cysteine residue not involved in maintaining the proper conformation of the antibody variant also may be
50

5 substituted, generally with serine, to improve the oxidative stability of
the molecule and prevent aberrant cross linking. Conversely, cysteine
bond(s) may be added to the antibody to improve its stability (particularly
10 where the antibody is an antibody fragment such as an Fv fragment).
5 Another type of amino acid variant has an altered glycosylation pattern.
This may be achieved by deleting one or more carbohydrate moieties found in
the antibody, and/or adding one or more glycosylation sites that are not
15 present in the antibody. Glycosylation of antibodies is typically either
N-linked or O-linked. N-linked refers to the attachment of the
10 carbohydrate moiety to the side chain of an asparagine residue. The
tripeptide sequences asparagine-X-serine and asparagine-X-threonine, where
20 X is any amino acid except proline, are the recognition sequences for
enzymatic attachment of the carbohydrate moiety to the asparagine side
chain. Thus, the presence of either of these tripeptide sequences in a
15 polypeptide creates a potential glycosylation site. O-linked glycosylation
refers to the attachment of one of the sugars N-acetylgalactosamine,
25 galactose, or xylose to a hydroxyamino acid, most commonly serine or
threonine, although 5-hydroxyproline or 5-hydroxylysine may also be used.
Addition of glycosylation sites to the antibody is conveniently
20 accomplished by altering the amino acid sequence such that it contains one
or more of the above-described tripeptide sequences (for N-linked
glycosylation sites). The alteration may also be made by the addition of,
or substitution by, one or more serine or threonine residues to the
30 sequence of the original antibody (for O-linked glycosylation sites).

35 25 Techniques for producing antibodies, which may be the parent antibody
and therefore require modification according to the techniques elaborated
herein, follow:

40 **A. Antibody Preparation**

(i) Antigen preparation

30 Soluble antigens or fragments thereof, optionally conjugated to other
molecules, can be used as immunogens for generating antibodies. For
transmembrane molecules, such as receptors, fragments of these (e.g. the
45 extracellular domain of a receptor) can be used as the immunogen.
Alternatively, cells expressing the transmembrane molecule can be used as
35 the immunogen. Such cells can be derived from a natural source (e.g.
cancer cell lines) or may be cells which have been transformed by
50 recombinant techniques to express the transmembrane molecule. Other

antigens and forms thereof useful for preparing antibodies will be apparent to those in the art.

(ii) *Polyclonal antibodies*

Polyclonal antibodies are preferably raised in animals by multiple subcutaneous (sc) or intraperitoneal (ip) injections of the relevant antigen and an adjuvant. It may be useful to conjugate the relevant antigen to a protein that is immunogenic in the species to be immunized, e.g., keyhole limpet hemocyanin, serum albumin, bovine thyroglobulin, or soybean trypsin inhibitor using a bifunctional or derivatizing agent, for example, maleimidobenzoyl sulfosuccinimide ester (conjugation through cysteine residues), N-hydroxysuccinimide (through lysine residues), glutaraldehyde, succinic anhydride, SOCl_2 , or $\text{R}^1\text{N}=\text{C}=\text{NR}$, where R and R^1 are different alkyl groups.

Animals are immunized against the antigen, immunogenic conjugates, or derivatives by combining, e.g., 100 μg or 5 μg of the protein or conjugate (for rabbits or mice, respectively) with 3 volumes of Freund's complete adjuvant and injecting the solution intradermally at multiple sites. One month later the animals are boosted with 1/5 to 1/10 the original amount of peptide or conjugate in Freund's complete adjuvant by subcutaneous injection at multiple sites. Seven to 14 days later the animals are bled and the serum is assayed for antibody titer. Animals are boosted until the titer plateaus. Preferably, the animal is boosted with the conjugate of the same antigen, but conjugated to a different protein and/or through a different cross-linking reagent. Conjugates also can be made in recombinant cell culture as protein fusions. Also, aggregating agents such as alum are suitably used to enhance the immune response.

(iii) *Monoclonal antibodies*

Monoclonal antibodies may be made using the hybridoma method first described by Kohler *et al.*, *Nature*, 256:495 (1975), or may be made by recombinant DNA methods (U.S. Patent No. 4,816,567).

In the hybridoma method, a mouse or other appropriate host animal, such as a hamster or macaque monkey, is immunized as hereinabove described to elicit lymphocytes that produce or are capable of producing antibodies that will specifically bind to the protein used for immunization. Alternatively, lymphocytes may be immunized *in vitro*. Lymphocytes then are fused with myeloma cells using a suitable fusing agent, such as polyethylene glycol, to form a hybridoma cell (Goding, *Monoclonal*

Antibodies: Principles and Practice, pp.59-103 (Academic Press, 1986)).

The hybridoma cells thus prepared are seeded and grown in a suitable culture medium that preferably contains one or more substances that inhibit the growth or survival of the unfused, parental myeloma cells. For example, if the parental myeloma cells lack the enzyme hypoxanthine guanine phosphoribosyl transferase (HGPRT or HPRT), the culture medium for the hybridomas typically will include hypoxanthine, aminopterin, and thymidine (HAT medium), which substances prevent the growth of HGPRT-deficient cells.

Preferred myeloma cells are those that fuse efficiently, support stable high-level production of antibody by the selected antibody-producing cells, and are sensitive to a medium such as HAT medium. Among these, preferred myeloma cell lines are murine myeloma lines, such as those derived from MOPC-21 and MPC-11 mouse tumors available from the Salk Institute Cell Distribution Center, San Diego, California USA, and SP-2 or X63-Ag8-653 cells available from the American Type Culture Collection, Rockville, Maryland USA. Human myeloma and mouse-human heteromyeloma cell lines also have been described for the production of human monoclonal antibodies (Kozbor, J. Immunol., 133:3001 (1984); Brodeur et al., Monoclonal Antibody Production Techniques and Applications, pp. 51-63 (Marcel Dekker, Inc., New York, 1987)).

Culture medium in which hybridoma cells are growing is assayed for production of monoclonal antibodies directed against the antigen. Preferably, the binding specificity of monoclonal antibodies produced by hybridoma cells is determined by immunoprecipitation or by an in vitro binding assay, such as radioimmunoassay (RIA) or enzyme-linked immunoabsorbent assay (ELISA).

After hybridoma cells are identified that produce antibodies of the desired specificity, affinity, and/or activity, the clones may be subcloned by limiting dilution procedures and grown by standard methods (Coding, Monoclonal Antibodies: Principles and Practice, pp.59-103 (Academic Press, 1986)). Suitable culture media for this purpose include, for example, D-MEM or RPMI-1640 medium. In addition, the hybridoma cells may be grown in vivo as ascites tumors in an animal.

The monoclonal antibodies secreted by the subclones are suitably separated from the culture medium, ascites fluid, or serum by conventional immunoglobulin purification procedures such as, for example, protein A-Sepharose, hydroxylapatite chromatography, gel electrophoresis, dialysis,

5 or affinity chromatography.

10 DNA encoding the monoclonal antibodies is readily isolated and sequenced using conventional procedures (e.g., by using oligonucleotide probes that are capable of binding specifically to genes encoding the heavy
5 and light chains of the monoclonal antibodies). The hybridoma cells serve as a preferred source of such DNA. Once isolated, the DNA may be placed into expression vectors, which are then transfected into host cells such as
15 *E. coli* cells, simian COS cells, Chinese Hamster Ovary (CHO) cells, or myeloma cells that do not otherwise produce immunoglobulin protein, to obtain the synthesis of monoclonal antibodies in the recombinant host cells. Recombinant production of antibodies will be described in more
20 detail below.

In a further embodiment, antibodies or antibody fragments can be isolated from antibody phage libraries generated using the techniques
15 described in McCafferty et al., *Nature*, 348:552-554 (1990). Clackson et al., *Nature*, 352:624-628 (1991) and Marks et al., *J. Mol. Biol.*, 222:581-597 (1991) describe the isolation of murine and human antibodies, respectively, using phage libraries. Subsequent publications describe the
25 production of high affinity (nM range) human antibodies by chain shuffling (Marks et al., *Bio/Technology*, 10:779-783 (1992)), as well as combinatorial infection and *in vivo* recombination as a strategy for constructing very large phage libraries (Waterhouse et al., *Nuc. Acids. Res.*, 21:2265-2266 (1993)). Thus, these techniques are viable alternatives to traditional
30 monoclonal antibody hybridoma techniques for isolation of monoclonal antibodies.

35 The DNA also may be modified, for example, by substituting the coding sequence for human heavy- and light-chain constant domains in place of the homologous murine sequences (U.S. Patent No. 4,816,567; Morrison, et al.,
40 *Proc. Natl Acad. Sci. USA*, 81:6851 (1984)), or by covalently joining to the immunoglobulin coding sequence all or part of the coding sequence for a non-immunoglobulin polypeptide.
30

45 Typically such non-immunoglobulin polypeptides are substituted for the constant domains of an antibody, or they are substituted for the variable domains of one antigen-combining site of an antibody to create a
35 chimeric bivalent antibody comprising one antigen-combining site having specificity for an antigen and another antigen-combining site having
50 specificity for a different antigen.

(iv) *Humanized and human antibodies*

A humanized antibody has one or more amino acid residues introduced into it from a source which is non-human. These non-human amino acid residues are often referred to as "import" residues, which are typically taken from an "import" variable domain. Humanization can be essentially performed following the method of Winter and co-workers (Jones et al., *Nature*, 321:522-525 (1986); Riechmann et al., *Nature*, 332:323-327 (1988); Verhoeven et al., *Science*, 239:1534-1536 (1988)), by substituting rodent CDRs or CDR sequences for the corresponding sequences of a human antibody. Accordingly, such "humanized" antibodies are chimeric antibodies (U.S. Patent No. 4,816,567) wherein substantially less than an intact human variable domain has been substituted by the corresponding sequence from a non-human species. In practice, humanized antibodies are typically human antibodies in which some CDR residues and possibly some FR residues are substituted by residues from analogous sites in rodent antibodies.

The choice of human variable domains, both light and heavy, to be used in making the humanized antibodies is very important to reduce antigenicity. According to the so-called "best-fit" method, the sequence of the variable domain of a rodent antibody is screened against the entire library of known human variable domain sequences. The human sequence which is closest to that of the rodent is then accepted as the human FR for the humanized antibody (Sims et al., *J. Immunol.*, 151:2296 (1993); Chothia et al., *J. Mol. Biol.*, 196:901 (1987)). Another method uses a particular FR derived from the consensus sequence of all human antibodies of a particular subgroup of light or heavy chains. The same FR may be used for several different humanized antibodies (Carter et al., *Proc. Natl. Acad. Sci. USA*, 89:4285 (1992); Presta et al., *J. Immunol.*, 151:2623 (1993)).

It is further important that antibodies be humanized with retention of high affinity for the antigen and other favorable biological properties. To achieve this goal, according to a preferred method, humanized antibodies are prepared by a process of analysis of the parental sequences and various conceptual humanized products using three-dimensional models of the parental and humanized sequences. Three-dimensional immunoglobulin models are commonly available and are familiar to those skilled in the art. Computer programs are available which illustrate and display probable three-dimensional conformational structures of selected candidate immunoglobulin sequences. Inspection of these displays permits analysis of

the likely role of the residues in the functioning of the candidate immunoglobulin sequence, i.e., the analysis of residues that influence the ability of the candidate immunoglobulin to bind its antigen. In this way, FR residues can be selected and combined from the recipient and import sequences so that the desired antibody characteristic, such as increased affinity for the target antigen(s), is achieved. In general, the CDR residues are directly and most substantially involved in influencing antigen binding.

Alternatively, it is now possible to produce transgenic animals (e.g., mice) that are capable, upon immunization, of producing a full repertoire of human antibodies in the absence of endogenous immunoglobulin production. For example, it has been described that the homozygous deletion of the antibody heavy-chain joining region (J_H) gene in chimeric and germ-line mutant mice results in complete inhibition of endogenous antibody production. Transfer of the human germ-line immunoglobulin gene array in such germ-line mutant mice will result in the production of human antibodies upon antigen challenge. See, e.g., Jakobovits et al., *Proc. Natl. Acad. Sci. USA*, 90:2551 (1993); Jakobovits et al., *Nature*, 362:255-258 (1993); Bruggermann et al., *Year in Immuno.*, 7:33 (1993); and Duchosal et al. *Nature* 355:258 (1992). Human antibodies can also be derived from phage-display libraries (Hoogenboom et al., *J. Mol. Biol.*, 227:381 (1991); Marks et al., *J. Mol. Biol.*, 222:581-597 (1991); Vaughan et al. *Nature Biotech* 14:309 (1996)).

(v) Antibody fragments

Various techniques have been developed for the production of antibody fragments. Traditionally, these fragments were derived via proteolytic digestion of intact antibodies (see, e.g., Morimoto et al., *Journal of Biochemical and Biophysical Methods* 24:107-117 (1992) and Brennan et al., *Science*, 229:81 (1985)). However, these fragments can now be produced directly by recombinant host cells. For example, the antibody fragments can be isolated from the antibody phage libraries discussed above. Alternatively, Fab'-SH fragments can be directly recovered from *E. coli* and chemically coupled to form F(ab')₂ fragments (Carter et al., *Bio/Technology* 10:163-167 (1992)). According to another approach, F(ab')₂ fragments can be isolated directly from recombinant host cell culture. Other techniques for the production of antibody fragments will be apparent to the skilled practitioner. In other embodiments, the antibody of choice is a single

chain Fv fragment (scFv). See WO 93/16185.

(vi) *Multispecific antibodies*

Multispecific antibodies have binding specificities for at least two different antigens. While such molecules normally will only bind two antigens (i.e. bispecific antibodies, BsAbs), antibodies with additional specificities such as trispecific antibodies are encompassed by this expression when used herein. Examples of BsAbs include those with one arm directed against a tumor cell antigen and the other arm directed against a cytotoxic trigger molecule such as anti-FcγRI/anti-CD15, anti-p185^{HER2}/FcγRIII (CD16), anti-CD3/anti-malignant B-cell (1D10), anti-CD3/anti-p185^{HER2}, anti-CD3/anti-p97, anti-CD3/anti-renal cell carcinoma, anti-CD3/anti-OVCAR-3, anti-CD3/L-D1 (anti-colon carcinoma), anti-CD3/anti-melanocyte stimulating hormone analog, anti-EGF receptor/anti-CD3, anti-CD3/anti-CAM11, anti-CD3/anti-CD19, anti-CD3/MoV18, anti-neural cell adhesion molecule (NCAM)/anti-CD3, anti-folate binding protein (FBP)/anti-CD3, anti-pan carcinoma associated antigen (AMOC-31)/anti-CD3; BsAbs with one arm which binds specifically to a tumor antigen and one arm which binds to a toxin such as anti-saporin/anti-Id-1, anti-CD22/anti-saporin, anti-CD7/anti-saporin, anti-CD38/anti-saporin, anti-CEA/anti-ricin A chain, anti-CEA/anti-vinca alkaloid; BsAbs for converting enzyme activated prodrugs such as anti-CD30/anti-alkaline phosphatase (which catalyzes conversion of mitomycin phosphate prodrug to mitomycin alcohol); BsAbs which can be used as fibrinolytic agents such as anti-fibrin/anti-tissue plasminogen activator (tPA), anti-fibrin/anti-urokinase-type plasminogen activator (uPA); BsAbs for targeting immune complexes to cell surface receptors such as anti-low density lipoprotein (LDL)/anti-Fc receptor (e.g. FcγRI, FcγRII or FcγRIII); BsAbs for use in therapy of infectious diseases such as anti-CD3/anti-herpes simplex virus (HSV), anti-T-cell receptor:CD3 complex/anti-influenza, anti-FcγR/anti-HIV; BsAbs for tumor detection in vitro or in vivo such as anti-CEA/anti-EOTUBE, anti-CEA/anti-DPTA, anti-p185^{HER2}/anti-hapten; BsAbs as vaccine adjuvants; and BsAbs as diagnostic tools such as anti-rabbit IgG/anti-ferritin, anti-horse radish peroxidase (HRP)/anti-hormone, anti-somatostatin/anti-substance P, anti-HRP/anti-FITC. Examples of trispecific antibodies include anti-CD3/anti-CD4/anti-CD37, anti-CD3/anti-CD5/anti-CD37 and anti-CD3/anti-CD8/anti-CD37. Bispecific antibodies can be prepared as full length antibodies or antibody fragments (e.g. F(ab')₂ bispecific antibodies).

5

Methods for making bispecific antibodies are known in the art. Traditional production of full length bispecific antibodies is based on the coexpression of two immunoglobulin heavy chain-light chain pairs, where the two chains have different specificities (Millstein et al., *Nature*, 305:537-539 (1983)). Because of the random assortment of immunoglobulin heavy and light chains, these hybridomas (quadromas) produce a potential mixture of 10 different antibody molecules, of which only one has the correct bispecific structure. Purification of the correct molecule, which is usually done by affinity chromatography steps, is rather cumbersome, and the product yields are low. Similar procedures are disclosed in WO 93/08829, and in Traunecker et al., *EMBO J.*, 10:3655-3659 (1991).

10

5

15

10

20

According to a different approach, antibody variable domains with the desired binding specificities (antibody-antigen combining sites) are fused to immunoglobulin constant domain sequences. The fusion preferably is with an immunoglobulin heavy chain constant domain, comprising at least part of the hinge, CH2, and CH3 regions. It is preferred to have the first heavy-chain constant region (CH1) containing the site necessary for light chain binding, present in at least one of the fusions. DNAs encoding the immunoglobulin heavy chain fusions and, if desired, the immunoglobulin light chain, are inserted into separate expression vectors, and are co-transfected into a suitable host organism. This provides for great flexibility in adjusting the mutual proportions of the three polypeptide fragments in embodiments when unequal ratios of the three polypeptide chains used in the construction provide the optimum yields. It is, however, possible to insert the coding sequences for two or all three polypeptide chains in one expression vector when the expression of at least two polypeptide chains in equal ratios results in high yields or when the ratios are of no particular significance.

25

30

35

25

40

30

45

35

50

In a preferred embodiment of this approach, the bispecific antibodies are composed of a hybrid immunoglobulin heavy chain with a first binding specificity in one arm, and a hybrid immunoglobulin heavy chain-light chain pair (providing a second binding specificity) in the other arm. It was found that this asymmetric structure facilitates the separation of the desired bispecific compound from unwanted immunoglobulin chain combinations, as the presence of an immunoglobulin light chain in only one half of the bispecific molecule provides for a facile way of separation. This approach is disclosed in WO 94/04690. For further details of

55

generating bispecific antibodies see, for example, Suresh et al., *Methods in Enzymology*, 121:210 (1986).

According to another approach described in W096/27011, the interface between a pair of antibody molecules can be engineered to maximize the percentage of heterodimers which are recovered from recombinant cell culture. The preferred interface comprises at least a part of the C_H3 domain of an antibody constant domain. In this method, one or more small amino acid side chains from the interface of the first antibody molecule are replaced with larger side chains (e.g. tyrosine or tryptophan). Compensatory "cavities" of identical or similar size to the large side chain(s) are created on the interface of the second antibody molecule by replacing large amino acid side chains with smaller ones (e.g. alanine or threonine). This provides a mechanism for increasing the yield of the heterodimer over other unwanted end-products such as homodimers.

Bispecific antibodies include cross-linked or "heteroconjugate" antibodies. For example, one of the antibodies in the heteroconjugate can be coupled to avidin, the other to biotin. Such antibodies have, for example, been proposed to target immune system cells to unwanted cells (US Patent No. 4,676,980), and for treatment of HIV infection (WO 91/00360, WO 92/200373, and EP 03089). Heteroconjugate antibodies may be made using any convenient cross-linking methods. Suitable cross-linking agents are well known in the art, and are disclosed in US Patent No. 4,676,980, along with a number of cross-linking techniques.

Techniques for generating bispecific antibodies from antibody fragments have also been described in the literature. For example, bispecific antibodies can be prepared using chemical linkage. Brennan et al., *Science*, 229: 81 (1985) describe a procedure wherein intact antibodies are proteolytically cleaved to generate F(ab')₂ fragments. These fragments are reduced in the presence of the dithiol complexing agent sodium arsenite to stabilize vicinal dithiols and prevent intermolecular disulfide formation. The Fab' fragments generated are then converted to thionitrobenzoate (TNB) derivatives. One of the Fab'-TNB derivatives is then reconverted to the Fab'-thiol by reduction with mercaptoethylamine and is mixed with an equimolar amount of the other Fab'-TNB derivative to form the bispecific antibody. The bispecific antibodies produced can be used as agents for the selective immobilization of enzymes.

Recent progress has facilitated the direct recovery of Fab'-SH

5

fragments from *E. coli*, which can be chemically coupled to form bispecific antibodies. Shalaby et al., *J. Exp. Med.*, 175: 217-225 (1992) describe the production of a fully humanized bispecific antibody F(ab')₂ molecule. Each Fab' fragment was separately secreted from *E. coli* and subjected to directed chemical coupling *in vitro* to form the bispecific antibody. The bispecific antibody thus formed was able to bind to cells overexpressing the ErbB2 receptor and normal human T cells, as well as trigger the lytic activity of human cytotoxic lymphocytes against human breast tumor targets.

10

5

15

Various techniques for making and isolating bispecific antibody fragments directly from recombinant cell culture have also been described. For example, bispecific antibodies have been produced using leucine zippers. Kostelny et al., *J. Immunol.*, 148(5):1547-1553 (1992). The leucine zipper peptides from the Fos and Jun proteins were linked to the Fab' portions of two different antibodies by gene fusion. The antibody homodimers were reduced at the hinge region to form monomers and then re-oxidized to form the antibody heterodimers. This method can also be utilized for the production of antibody homodimers. The "diabody" technology described by Hollinger et al., *Proc. Natl. Acad. Sci. USA*, 90:6444-6448 (1993) has provided an alternative mechanism for making bispecific antibody fragments. The fragments comprise a heavy-chain variable domain (V_H) connected to a light-chain variable domain (V_L) by a linker which is too short to allow pairing between the two domains on the same chain. Accordingly, the V_H and V_L domains of one fragment are forced to pair with the complementary V_L and V_H domains of another fragment, thereby forming two antigen-binding sites. Another strategy for making bispecific antibody fragments by the use of single-chain Fv (sFv) dimers has also been reported. See Gruber et al., *J. Immunol.*, 152:5368 (1994).

10

20

25

30

35

25

40

30

(vii) Effector function engineering

It may be desirable to modify the antibody of the invention with respect to effector function, so as to enhance the effectiveness of the antibody in treating cancer, for example. For example cysteine residue(s) may be introduced in the Fc region, thereby allowing interchain disulfide bond formation in this region. The homodimeric antibody thus generated may have improved internalization capability and/or increased complement-

45

35

50

55

mediated cell killing and antibody-dependent cellular cytotoxicity (ADCC). See Caron et al., *J. Exp Med.* 176:1191-1195 (1992) and Shopes, B. J. *Immunol.* 148:2918-2922 (1992). Homodimeric antibodies with enhanced anti-tumor activity may also be prepared using heterobifunctional cross-linkers as described in Wolff et al. *Cancer Research* 53:2560-2565 (1993). Alternatively, an antibody can be engineered which has dual Fc regions and may thereby have enhanced complement lysis and ADCC capabilities. See Stevenson et al. *Anti-Cancer Drug Design* 3:219-230 (1989).

(viii) *Immunoconjugates*

The invention also pertains to immunoconjugates comprising the antibody described herein conjugated to a cytotoxic agent such as a chemotherapeutic agent, toxin (e.g. an enzymatically active toxin of bacterial, fungal, plant or animal origin, or fragments thereof), or a radioactive isotope (i.e., a radioconjugate).

Chemotherapeutic agents useful in the generation of such immunoconjugates have been described above. Enzymatically active toxins and fragments thereof which can be used include diphtheria A chain, nonbinding active fragments of diphtheria toxin, exotoxin A chain (from *Pseudomonas aeruginosa*), ricin A chain, abrin A chain, modeccin A chain, alpha-sarcin, *Aleurites fordii* proteins, dianthin proteins, *Phytolacca americana* proteins (PAPI, PAPII, and PAP-S), momordica charantia inhibitor, curcin, crotin, sapaonaria officinalis inhibitor, gelonin, mitogellin, restrictocin, phenomycin, enomycin and the tricothecenes. A variety of radionuclides are available for the production of radioconjugate antibodies. Examples include ^{212}Bi , ^{131}I , ^{131}In , ^{90}Y and ^{186}Re .

Conjugates of the antibody and cytotoxic agent are made using a variety of bifunctional protein coupling agents such as N-succinimidyl-3-(2-pyridyldithiol) propionate (SPDP), iminothiolane (IT), bifunctional derivatives of imidoesters (such as dimethyl adipimidate HCL), active esters (such as disuccinimidyl suberate), aldehydes (such as glutaraldehyde), bis-azido compounds (such as bis (p-azidobenzoyl) hexanediamine), bis-diazonium derivatives (such as bis-(p-diazoniumbenzoyl)-ethylenediamine), diisocyanates (such as tolyene 2,6-diisocyanate), and bis-active fluorine compounds (such as 1,5-difluoro-2,4-dinitrobenzene). For example, a ricin immunotoxin can be prepared as described in Vitetta et al. *Science* 238: 1098 (1987). Carbon-14-labeled 1-isothiocyanatobenzyl-3-methyldiethylene triaminepentaacetic acid (MX-DTPA)

5 is an exemplary chelating agent for conjugation of radionucleotide to the antibody. See WO94/11026.

10 In another embodiment, the antibody may be conjugated to a "receptor" (such streptavidin) for utilization in tumor pretargeting wherein the
5 antibody-receptor conjugate is administered to the patient, followed by removal of unbound conjugate from the circulation using a clearing agent and then administration of a "ligand" (e.g. avidin) which is conjugated to
15 a cytotoxic agent (e.g. a radionucleotide).

(ix) Immunoliposomes.

10 The antibody variants disclosed herein may also be formulated as immunoliposomes. Liposomes containing the antibody are prepared by
20 methods known in the art, such as described in Epstein et al., Proc. Natl. Acad. Sci. USA, 82:3688 (1985); Hwang et al., Proc. Natl. Acad. Sci. USA, 77:4030 (1980); and U.S. Pat. Nos. 4,485,045 and 4,544,545. Liposomes with
15 enhanced circulation time are disclosed in U.S. Patent No. 5,013,556.

25 Particularly useful liposomes can be generated by the reverse phase evaporation method with a lipid composition comprising phosphatidylcholine, cholesterol and PEG-derivatized phosphatidylethanolamine (PEG-PE). Liposomes are extruded through filters of defined pore size to yield
30 liposomes with the desired diameter. Fab' fragments of the antibody of the present invention can be conjugated to the liposomes as described in Martin et al. J. Biol. Chem. 257: 285-288 (1982) via a disulfide interchange reaction. A chemotherapeutic agent (such as Doxorubicin) is optionally
35 contained within the liposome. See Gabizon et al. J. National Cancer Inst. 81(19):1484 (1989)

(x) Antibody Dependent Enzyme Mediated Prodrug Therapy (ADEPT)

40 The antibody of the present invention may also be used in ADEPT by conjugating the antibody to a prodrug-activating enzyme which converts a prodrug (e.g. a peptidyl chemotherapeutic agent, see WO81/01145) to an
30 active anti-cancer drug. See, for example, WO 88/07378 and U.S. Patent No. 4,975,278.

45 The enzyme component of the immunoconjugate useful for ADEPT includes any enzyme capable of acting on a prodrug in such a way so as to convert it into its more active, cytotoxic form.

35 Enzymes that are useful in the method of this invention include, but are not limited to, alkaline phosphatase useful for converting phosphate-containing prodrugs into free drugs; arylsulfatase useful for converting
50

5 sulfate-containing prodrugs into free drugs; cytosine deaminase useful for
converting non-toxic 5-fluorocytosine into the anti-cancer drug, 5-
10 fluorouracil; proteases, such as serratia protease, thermolysin,
subtilisin, carboxypeptidases and cathepsins (such as cathepsins B and L),
5 that are useful for converting peptide-containing prodrugs into free drugs;
D-alanylcarboxypeptidases, useful for converting prodrugs that contain D-
15 amino acid substituents; carbohydrate-cleaving enzymes such as beta-
galactosidase and neuraminidase useful for converting glycosylated prodrugs
into free drugs; beta-lactamase useful for converting drugs derivatized
20 with beta-lactams into free drugs; and penicillin amidases, such as
penicillin V amidase or penicillin G amidase, useful for converting drugs
derivatized at their amine nitrogens with phenoxyacetyl or phenylacetyl
groups, respectively, into free drugs. Alternatively, antibodies with
enzymatic activity, also known in the art as "abzymes", can be used to
15 convert the prodrugs of the invention into free active drugs (see, e.g.,
Massey, *Nature* 328: 457-458 (1987)). Antibody-abzyme conjugates can be
25 prepared as described herein for delivery of the abzyme to a tumor cell
population.

The enzymes of this invention can be covalently bound to the antibody
30 variant by techniques well known in the art such as the use of the
heterobifunctional crosslinking reagents discussed above. Alternatively,
fusion proteins comprising at least the antigen binding region of an
antibody of the invention linked to at least a functionally active portion
of an enzyme of the invention can be constructed using recombinant DNA
35 techniques well known in the art (see, e.g., Neuberger et al., *Nature*, 312:
25 604-608 (1984)).

(xi) Antibody-salvage receptor binding epitope fusions.

In certain embodiments of the invention, it may be desirable to use
40 an antibody fragment, rather than an intact antibody, to increase tumor
penetration, for example. In this case, it may be desirable to modify the
30 antibody fragment in order to increase its serum half life. This may be
achieved, for example, by incorporation of a salvage receptor binding
45 epitope into the antibody fragment (e.g. by mutation of the appropriate
region in the antibody fragment or by incorporating the epitope into a
peptide tag that is then fused to the antibody fragment at either end or in
35 the middle, e.g., by DNA or peptide synthesis).

The salvage receptor binding epitope preferably constitutes a region

5 wherein any one or more amino acid residues from one or two loops of a Fc domain are transferred to an analogous position of the antibody fragment. Even more preferably, three or more residues from one or two loops of the
10 Fc domain are transferred. Still more preferred, the epitope is taken from the CH2 domain of the Fc region (e.g., of an IgG) and transferred to the
5 CH1, CH3, or V_H region, or more than one such region, of the antibody. Alternatively, the epitope is taken from the CH2 domain of the Fc region and transferred to the C_L region or V_L region, or both, of the antibody
15 fragment. See, e.g., US Patent 5,739,277, issued April 14, 1998.

10 (xii) Covalent modifications

Covalent modifications of the antibody are included within the scope
20 of this invention. They may be made by chemical synthesis or by enzymatic or chemical cleavage of the antibody, if applicable. Other types of covalent modifications of the antibody are introduced into the molecule by
15 reacting targeted amino acid residues of the antibody with an organic derivatizing agent that is capable of reacting with selected side chains or
25 the N- or C-terminal residues.

Removal of any carbohydrate moieties present on the antibody may be accomplished chemically or enzymatically. Chemical deglycosylation
30 requires exposure of the antibody to the compound trifluoromethanesulfonic acid, or an equivalent compound. This treatment results in the cleavage of most or all sugars except the linking sugar (N-acetylglucosamine or N-acetylgalactosamine), while leaving the antibody intact. Chemical
35 deglycosylation is described by Hakimuddin, et al. *Arch. Biochem. Biophys.* 259:52 (1987) and by Edge et al. *Anal. Biochem.*, 118:131 (1981). Enzymatic cleavage of carbohydrate moieties on antibodies can be achieved by the use of a variety of endo- and exo-glycosidases as described by Thotakura et al.
40 *Meth. Enzymol.* 138:350 (1987).

Another type of covalent modification of the antibody comprises
40 linking the antibody to one of a variety of nonproteinaceous polymers, e.g., polyethylene glycol, polypropylene glycol, or polyoxyalkylenes, in the manner set forth in U.S. Patent Nos. 4,640,835; 4,496,639; 4,301,144;
45 4,670,417; 4,791,192 or 4,179,337.

B. Vectors, Host Cells and Recombinant Methods

35 The invention also provides isolated nucleic acid encoding an antibody variant as disclosed herein, vectors and host cells comprising the nucleic acid, and recombinant techniques for the production of the antibody
50

variant.

For recombinant production of the antibody variant, the nucleic acid encoding it is isolated and inserted into a replicable vector for further cloning (amplification of the DNA) or for expression. DNA encoding the monoclonal antibody variant is readily isolated and sequenced using conventional procedures (e.g., by using oligonucleotide probes that are capable of binding specifically to genes encoding the heavy and light chains of the antibody variant). Many vectors are available. The vector components generally include, but are not limited to, one or more of the following: a signal sequence, an origin of replication, one or more marker genes, an enhancer element, a promoter, and a transcription termination sequence.

(i) *Signal sequence component*

The antibody variant of this invention may be produced recombinantly not only directly, but also as a fusion polypeptide with a heterologous polypeptide, which is preferably a signal sequence or other polypeptide having a specific cleavage site at the N-terminus of the mature protein or polypeptide. The heterologous signal sequence selected preferably is one that is recognized and processed (i.e., cleaved by a signal peptidase) by the host cell. For prokaryotic host cells that do not recognize and process the native antibody signal sequence, the signal sequence is substituted by a prokaryotic signal sequence selected, for example, from the group of the alkaline phosphatase, penicillinase, lpp, or heat-stable enterotoxin II leaders. For yeast secretion the native signal sequence may be substituted by, e.g., the yeast invertase leader, α -factor leader (including *Saccharomyces* and *Kluyveromyces* α -factor leaders), or acid phosphatase leader, the *C. albicans* glucoamylase leader, or the signal described in WO 90/13646. In mammalian cell expression, mammalian signal sequences as well as viral secretory leaders, for example, the herpes simplex gD signal, are available.

The DNA for such precursor region is ligated in reading frame to DNA encoding the antibody variant.

(ii) *Origin of replication component*

Both expression and cloning vectors contain a nucleic acid sequence that enables the vector to replicate in one or more selected host cells. Generally, in cloning vectors this sequence is one that enables the vector to replicate independently of the host chromosomal DNA, and includes

origins of replication or autonomously replicating sequences. Such sequences are well known for a variety of bacteria, yeast, and viruses. The origin of replication from the plasmid pBR322 is suitable for most Gram-negative bacteria, the 2μ plasmid origin is suitable for yeast, and various viral origins (SV40, polyoma, adenovirus, VSV or BPV) are useful for cloning vectors in mammalian cells. Generally, the origin of replication component is not needed for mammalian expression vectors (the SV40 origin may typically be used only because it contains the early promoter).

(iii) Selection gene component

Expression and cloning vectors may contain a selection gene, also termed a selectable marker. Typical selection genes encode proteins that (a) confer resistance to antibiotics or other toxins, e.g., ampicillin, neomycin, methotrexate, or tetracycline, (b) complement auxotrophic deficiencies, or (c) supply critical nutrients not available from complex media, e.g., the gene encoding D-alanine racemase for *Bacilli*.

One example of a selection scheme utilizes a drug to arrest growth of a host cell. Those cells that are successfully transformed with a heterologous gene produce a protein conferring drug resistance and thus survive the selection regimen. Examples of such dominant selection use the drugs neomycin, mycophenolic acid and hygromycin.

Another example of suitable selectable markers for mammalian cells are those that enable the identification of cells competent to take up the antibody nucleic acid, such as DHFR, thymidine kinase, metallothionein-I and -II, preferably primate metallothionein genes, adenosine deaminase, ornithine decarboxylase, etc.

For example, cells transformed with the DHFR selection gene are first identified by culturing all of the transformants in a culture medium that contains methotrexate (Mtx), a competitive antagonist of DHFR. An appropriate host cell when wild-type DHFR is employed is the Chinese hamster ovary (CHO) cell line deficient in DHFR activity.

Alternatively, host cells (particularly wild-type hosts that contain endogenous DHFR) transformed or co-transformed with DNA sequences encoding antibody, wild-type DHFR protein, and another selectable marker such as aminoglycoside 3'-phosphotransferase (APH) can be selected by cell growth in medium containing a selection agent for the selectable marker such as an aminoglycosidic antibiotic, e.g., kanamycin, neomycin, or G418. See U.S.

5

Patent No. 4,965,199.

10

A suitable selection gene for use in yeast is the *trp1* gene present in the yeast plasmid YRp7 (Stinchcomb et al., *Nature*, 282:39 (1979)). The *trp1* gene provides a selection marker for a mutant strain of yeast lacking the ability to grow in tryptophan, for example, ATCC No. 44076 or PEP4-1. Jones, *Genetics*, 85:12 (1977). The presence of the *trp1* lesion in the yeast host cell genome then provides an effective environment for detecting transformation by growth in the absence of tryptophan. Similarly, *Leu2*-deficient yeast strains (ATCC 20,622 or 38,626) are complemented by known plasmids bearing the *Leu2* gene.

15

In addition, vectors derived from the 1.6 μ m circular plasmid pKD1 can be used for transformation of *Kluyveromyces* yeasts. Alternatively, an expression system for large-scale production of recombinant calf chymosin was reported for *K. lactis*. Van den Berg, *Bio/Technology*, 8:135 (1990). Stable multi-copy expression vectors for secretion of mature recombinant human serum albumin by industrial strains of *Kluyveromyces* have also been disclosed. Fleer et al., *Bio/Technology*, 9:968-975 (1991).

20

25

(iv) Promoter component

30

Expression and cloning vectors usually contain a promoter that is recognized by the host organism and is operably linked to the antibody nucleic acid. Promoters suitable for use with prokaryotic hosts include the *phoA* promoter, beta-lactamase and lactose promoter systems, alkaline phosphatase, a tryptophan (*trp*) promoter system, and hybrid promoters such as the *tac* promoter. However, other known bacterial promoters are suitable. Promoters for use in bacterial systems also will contain a Shine-Dalgarno (S.D.) sequence operably linked to the DNA encoding the antibody.

35

40

Promoter sequences are known for eukaryotes. Virtually all eukaryotic genes have an AT-rich region located approximately 25 to 30 bases upstream from the site where transcription is initiated. Another sequence found 70 to 80 bases upstream from the start of transcription of many genes is a CNCAAT region where N may be any nucleotide. At the 3' end of most eukaryotic genes is an AATAAA sequence that may be the signal for addition of the poly A tail to the 3' end of the coding sequence. All of these sequences are suitably inserted into eukaryotic expression vectors.

45

Examples of suitable promoting sequences for use with yeast hosts include the promoters for 3-phosphoglycerate kinase or other glycolytic

50

55

5 enzymes, such as enolase, glyceraldehyde-3 phosphate dehydrogenase, hexokinase, pyruvate decarboxylase, phosphofructokinase, glucose-6-phosphate isomerase, 3-phosphoglycerate mutase, pyruvate kinase,
10 triosephosphate isomerase, phosphoglucose isomerase, and glucokinase.

5 Other yeast promoters, which are inducible promoters having the additional advantage of transcription controlled by growth conditions, are the promoter regions for alcohol dehydrogenase 2, isocytochrome C, acid
15 phosphatase, degradative enzymes associated with nitrogen metabolism, metallothionein, glyceraldehyde-3-phosphate dehydrogenase, and enzymes
20 responsible for maltose and galactose utilization. Suitable vectors and promoters for use in yeast expression are further described in EP 73,657. Yeast enhancers also are advantageously used with yeast promoters.

Antibody transcription from vectors in mammalian host cells is controlled, for example, by promoters obtained from the genomes of viruses
15 such as polyoma virus, fowlpox virus, adenovirus (such as Adenovirus 2), bovine papilloma virus, avian sarcoma virus, cytomegalovirus, a retrovirus,
25 hepatitis-B virus and most preferably Simian Virus 40 (SV40), from heterologous mammalian promoters, e.g., the actin promoter or an immunoglobulin promoter, from heat-shock promoters, provided such promoters
30 are compatible with the host cell systems.

The early and late promoters of the SV40 virus are conveniently obtained as an SV40 restriction fragment that also contains the SV40 viral origin of replication. The immediate early promoter of the human
35 cytomegalovirus is conveniently obtained as a HindIII E restriction
25 fragment. A system for expressing DNA in mammalian hosts using the bovine papilloma virus as a vector is disclosed in U.S. Patent No. 4,419,446. A modification of this system is described in U.S. Patent No. 4,601,978. Alternatively, the rous sarcoma virus long terminal repeat can be used as the promoter.

30 (v) *Enhancer element component*

Transcription of a DNA encoding the antibody of this invention by higher eukaryotes is often increased by inserting an enhancer sequence into
45 the vector. Many enhancer sequences are now known from mammalian genes (globin, elastase, albumin, alpha-fetoprotein, and insulin). Typically,
35 however, one will use an enhancer from a eukaryotic cell virus. Examples include the SV40 enhancer on the late side of the replication origin (bp
50 100-270), the cytomegalovirus early promoter enhancer, the polyoma enhancer

5 on the late side of the replication origin, and adenovirus enhancers. See
also Yaniv, *Nature* 297:17-18 (1982) on enhancing elements for activation of
eukaryotic promoters. The enhancer may be spliced into the vector at a
10 position 5' or 3' to the antibody-encoding sequence, but is preferably
5 located at a site 5' from the promoter.

(vi) *Transcription termination component*

15 Expression vectors used in eukaryotic host cells (yeast, fungi,
insect, plant, animal, human, or nucleated cells from other multicellular
organisms) will also contain sequences necessary for the termination of
10 transcription and for stabilizing the mRNA. Such sequences are commonly
available from the 5' and, occasionally 3', untranslated regions of
20 eukaryotic or viral DNAs or cDNAs. These regions contain nucleotide
segments transcribed as polyadenylated fragments in the untranslated
portion of the mRNA encoding the antibody. One useful transcription
15 termination component is the bovine growth hormone polyadenylation region.
See WO94/11026 and the expression vector disclosed therein.

25 (vii) *Selection and transformation of host cells*

Suitable host cells for cloning or expressing the DNA in the vectors
herein are the prokaryote, yeast, or higher eukaryote cells described
20 above. Suitable prokaryotes for this purpose include eubacteria, such as
Gram-negative or Gram-positive organisms, for example, Enterobacteriaceae
such as *Escherichia*, e.g., *E. coli*, *Enterobacter*, *Erwinia*, *Klebsiella*,
Proteus, *Salmonella*, e.g., *Salmonella typhimurium*, *Serratia*, e.g., *Serratia*
marcescans, and *Shigella*, as well as *Bacilli* such as *B. subtilis* and *B.*
35 *licheniformis* (e.g., *B. licheniformis* 41P disclosed in DD 266,710 published
12 April 1989), *Pseudomonas* such as *P. aeruginosa*, and *Streptomyces*. One
preferred *E. coli* cloning host is *E. coli* 294 (ATCC 31,446), although other
strains such as *E. coli* B, *E. coli* X1776 (ATCC 31,537), and *E. coli* W3110
40 (ATCC 27,325) are suitable. These examples are illustrative rather than
30 limiting.

In addition to prokaryotes, eukaryotic microbes such as filamentous
fungi or yeast are suitable cloning or expression hosts for antibody-
45 encoding vectors. *Saccharomyces cerevisiae*, or common baker's yeast, is
the most commonly used among lower eukaryotic host microorganisms.
35 However, a number of other genera, species, and strains are commonly
available and useful herein, such as *Schizosaccharomyces pombe*,
50 *Kluyveromyces* hosts such as, e.g., *K. lactis*, *K. fragilis* (ATCC 12,424), *K.*

5 *bulgaricus* (ATCC 15,045), *K. wickramii* (ATCC 24,178), *K. waltii* (ATCC
56,500), *K. drosophilum* (ATCC 36,906), *K. thermotolerans*, and *K.*
10 *marxianus*; *Yarrowia* (EP 402,226); *Pichia pastoris* (EP 183,070); *Candida*;
15 *Trichoderma reesia* (EP 244,234); *Neurospora crassa*; *Schwanniomyces* such as
5 *Schwanniomyces occidentalis*; and filamentous fungi such as, e.g.,
Neurospora, *Penicillium*, *Tolypocladium*, and *Aspergillus* hosts such as *A.*
nidulans and *A. niger*.

15 Suitable host cells for the expression of glycosylated antibody are
derived from multicellular organisms. Examples of invertebrate cells
20 include plant and insect cells. Numerous baculoviral strains and variants
and corresponding permissive insect host cells from hosts such as
30 *Spodoptera frugiperda* (caterpillar), *Aedes aegypti* (mosquito), *Aedes*
albopictus (mosquito), *Drosophila melanogaster* (fruitfly), and *Bombyx mori*
have been identified. A variety of viral strains for transfection are
15 publicly available, e.g., the L-1 variant of *Autographa californica* NPV and
the Bm-5 strain of *Bombyx mori* NPV, and such viruses may be used as the
25 virus herein according to the present invention, particularly for
transfection of *Spodoptera frugiperda* cells. Plant cell cultures of
cotton, corn, potato, soybean, petunia, tomato, and tobacco can also be
30 utilized as hosts.

 However, interest has been greatest in vertebrate cells, and
propagation of vertebrate cells in culture (tissue culture) has become a
routine procedure. Examples of useful mammalian host cell lines are monkey
35 kidney CV1 line transformed by SV40 (COS-7, ATCC CRL 1651); human embryonic
25 kidney line (293 or 293 cells subcloned for growth in suspension culture,
Graham et al., *J. Gen. Virol.* 36:59 (1977)); baby hamster kidney cells (BHK,
ATCC CCL 10); Chinese hamster ovary cells/-DHFR (CHO, Urlaub et al., *Proc.*
Natl. Acad. Sci. USA 77:4216 (1980)); mouse sertoli cells (TM4, Mather,
40 *Biol. Reprod.* 23:243-251 (1980)); monkey kidney cells (CV1 ATCC CCL 70);
30 African green monkey kidney cells (VERO-76, ATCC CRL-1587); human cervical
carcinoma cells (HELA, ATCC CCL 2); canine kidney cells (MDCK, ATCC CCL
34); buffalo rat liver cells (BRL 3A, ATCC CRL 1442); human lung cells
45 (W138, ATCC CCL 75); human liver cells (Hep G2, HB 8065); mouse mammary
tumor (MMT 060562, ATCC CCL51); TRI cells (Mather et al., *Annals N.Y. Acad.*
35 *Sci.* 383:44-68 (1982)); MRC 5 cells; FS4 cells; and a human hepatoma line
(Hep G2).

50 Host cells are transformed with the above-described expression or

cloning vectors for antibody production and cultured in conventional nutrient media modified as appropriate for inducing promoters, selecting transformants, or amplifying the genes encoding the desired sequences.

(viii) *Culturing the host cells*

The host cells used to produce the antibody variant of this invention may be cultured in a variety of media. Commercially available media such as Ham's F10 (Sigma), Minimal Essential Medium ((MEM), (Sigma), RPMI-1640 (Sigma), and Dulbecco's Modified Eagle's Medium ((DMEM), Sigma) are suitable for culturing the host cells. In addition, any of the media described in Ham et al., *Meth. Enz.* 58:44 (1979), Barnes et al., *Anal. Biochem.* 102:255 (1980), U.S. Pat. Nos. 4,767,704; 4,657,866; 4,927,762; 4,560,655; or 5,122,469; WO 90/03430; WO 87/00195; or U.S. Patent Re. 30,985 may be used as culture media for the host cells. Any of these media may be supplemented as necessary with hormones and/or other growth factors (such as insulin, transferrin, or epidermal growth factor), salts (such as sodium chloride, calcium, magnesium, and phosphate), buffers (such as HEPES), nucleotides (such as adenosine and thymidine), antibiotics (such as GENTAMYCIN[®]drug), trace elements (defined as inorganic compounds usually present at final concentrations in the micromolar range), and glucose or an equivalent energy source. Any other necessary supplements may also be included at appropriate concentrations that would be known to those skilled in the art. The culture conditions, such as temperature, pH, and the like, are those previously used with the host cell selected for expression, and will be apparent to the ordinarily skilled artisan.

(ix) *Antibody purification*

When using recombinant techniques, the antibody variant can be produced intracellularly, in the periplasmic space, or directly secreted into the medium. If the antibody variant is produced intracellularly, as a first step, the particulate debris, either host cells or lysed fragments, is removed, for example, by centrifugation or ultrafiltration. Carter et al., *Bio/Technology* 10:163-167 (1992) describe a procedure for isolating antibodies which are secreted to the periplasmic space of *E. coli*. Briefly, cell paste is thawed in the presence of sodium acetate (pH 3.5), EDTA, and phenylmethylsulfonylfluoride (PMSF) over about 30 min. Cell debris can be removed by centrifugation. Where the antibody variant is secreted into the medium, supernatants from such expression systems are generally first concentrated using a commercially available protein

concentration filter, for example, an Amicon or Millipore Pellicon ultrafiltration unit. A protease inhibitor such as PMSF may be included in any of the foregoing steps to inhibit proteolysis and antibiotics may be included to prevent the growth of adventitious contaminants.

The antibody composition prepared from the cells can be purified using, for example, hydroxylapatite chromatography, gel electrophoresis, dialysis, and affinity chromatography, with affinity chromatography being the preferred purification technique. The suitability of protein A as an affinity ligand depends on the species and isotype of any immunoglobulin Fc domain that is present in the antibody variant. Protein A can be used to purify antibodies that are based on human $\gamma 1$, $\gamma 2$, or $\gamma 4$ heavy chains (Lindmark et al., *J. Immunol. Meth.* 62:1-13 (1983)). Protein G is recommended for all mouse isotypes and for human $\gamma 3$ (Guss et al., *EMBO J.* 5:1567-1575 (1986)). The matrix to which the affinity ligand is attached is most often agarose, but other matrices are available. Mechanically stable matrices such as controlled pore glass or poly(styrene-divinyl)benzene allow for faster flow rates and shorter processing times than can be achieved with agarose. Where the antibody variant comprises a C_H3 domain, the Bakerbond ABXTM resin (J. T. Baker, Phillipsburg, NJ) is useful for purification. Other techniques for protein purification such as fractionation on an ion-exchange column, ethanol precipitation, Reverse Phase HPLC, chromatography on silica, chromatography on heparin SEPHAROSETM chromatography on an anion or cation exchange resin (such as a polyaspartic acid column), chromatofocusing, SDS-PAGE, and ammonium sulfate precipitation are also available depending on the antibody variant to be recovered.

C. Pharmaceutical Formulations

Therapeutic formulations of the antibody variant are prepared for storage by mixing the antibody variant having the desired degree of purity with optional physiologically acceptable carriers, excipients or stabilizers (*Remington's Pharmaceutical Sciences* 16th edition, Osol, A. Ed. (1980)), in the form of lyophilized formulations or aqueous solutions. Acceptable carriers, excipients, or stabilizers are nontoxic to recipients at the dosages and concentrations employed, and include buffers such as phosphate, citrate, and other organic acids; antioxidants including ascorbic acid and methionine; preservatives (such as octadecyldimethylbenzyl ammonium chloride; hexamethonium chloride;

5 benzalkonium chloride, benzethonium chloride; phenol, butyl or benzyl
alcohol; alkyl parabens such as methyl or propyl paraben; catechol;
10 resorcinol; cyclohexanol; 3-pentanol; and m-cresol); low molecular weight
(less than about 10 residues) polypeptide; proteins, such as serum albumin,
5 gelatin, or immunoglobulins; hydrophilic polymers such as
polyvinylpyrrolidone; amino acids such as glycine, glutamine, asparagine,
histidine, arginine, or lysine; monosaccharides, disaccharides, and other
15 carbohydrates including glucose, mannose, or dextrans; chelating agents
such as EDTA; sugars such as sucrose, mannitol, trehalose or sorbitol;
10 salt-forming counter-ions such as sodium; metal complexes (e.g., Zn-protein
complexes); and/or non-ionic surfactants such as TWEENTM, PLURONICSTM or
20 polyethylene glycol (PEG).

The formulation herein may also contain more than one active compound
as necessary for the particular indication being treated, preferably those
15 with complementary activities that do not adversely affect each other. For
example, it may be desirable to further provide an immunosuppressive agent.
25 Such molecules are suitably present in combination in amounts that are
effective for the purpose intended.

The active ingredients may also be entrapped in microcapsule
20 prepared, for example, by coacervation techniques or by interfacial
polymerization, for example, hydroxymethylcellulose or gelatin-microcapsule
and poly-(methylmethacrylate) microcapsule, respectively, in colloidal drug
delivery systems (for example, liposomes, albumin microspheres,
35 microemulsions, nano-particles and nanocapsules) or in macroemulsions.
25 Such techniques are disclosed in *Remington's Pharmaceutical Sciences* 16th
edition, Osol, A. Ed. (1980).

The formulations to be used for *in vivo* administration must be
sterile. This is readily accomplished by filtration through sterile
40 filtration membranes.

30 Sustained-release preparations may be prepared. Suitable examples of
sustained-release preparations include semipermeable matrices of solid
hydrophobic polymers containing the antibody variant, which matrices are in
the form of shaped articles, e.g., films, or microcapsule. Examples of
45 sustained-release matrices include polyesters, hydrogels (for example,
poly(2-hydroxyethyl-methacrylate), or poly(vinylalcohol)), polylactides
35 (U.S. Pat. No. 3,773,919), copolymers of L-glutamic acid and ethyl-L-
glutamate, non-degradable ethylene-vinyl acetate, degradable lactic acid-

glycolic acid copolymers such as the LUPRON DEPOT™ (injectable microspheres composed of lactic acid-glycolic acid copolymer and leuprolide acetate), and poly-D-(-)-3-hydroxybutyric acid. While polymers such as ethylene-vinyl acetate and lactic acid-glycolic acid enable release of molecules for over 100 days, certain hydrogels release proteins for shorter time periods. When encapsulated antibodies remain in the body for a long time, they may denature or aggregate as a result of exposure to moisture at 37°C, resulting in a loss of biological activity and possible changes in immunogenicity. Rational strategies can be devised for stabilization depending on the mechanism involved. For example, if the aggregation mechanism is discovered to be intermolecular S-S bond formation through thio-disulfide interchange, stabilization may be achieved by modifying sulfhydryl residues, lyophilizing from acidic solutions, controlling moisture content, using appropriate additives, and developing specific polymer matrix compositions.

D. Non-Therapeutic Uses for the Antibody Variant

The antibody variants of the invention may be used as affinity purification agents. In this process, the antibodies are immobilized on a solid phase such as a Sephadex resin or filter paper, using methods well known in the art. The immobilized antibody variant is contacted with a sample containing the antigen to be purified, and thereafter the support is washed with a suitable solvent that will remove substantially all the material in the sample except the antigen to be purified, which is bound to the immobilized antibody variant. Finally, the support is washed with another suitable solvent, such as glycine buffer, pH 5.0, that will release the antigen from the antibody variant.

The variant antibodies may also be useful in diagnostic assays, e.g., for detecting expression of an antigen of interest in specific cells, tissues, or serum.

For diagnostic applications, the antibody variant typically will be labeled with a detectable moiety. Numerous labels are available which can be generally grouped into the following categories:

(a) Radioisotopes, such as ^{35}S , ^{14}C , ^{125}I , ^3H , and ^{131}I . The antibody variant can be labeled with the radioisotope using the techniques described in *Current Protocols in Immunology*, Volumes 1 and 2, Coligan et al., Ed. Wiley-Interscience, New York, New York, Pubs. (1991) for example and radioactivity can be measured using scintillation counting.

(b) Fluorescent labels such as rare earth chelates (europium chelates) or fluorescein and its derivatives, rhodamine and its derivatives, dansyl, Lissamine, phycoerythrin and Texas Red are available. The fluorescent labels can be conjugated to the antibody variant using the techniques disclosed in *Current Protocols in Immunology*, supra, for example. Fluorescence can be quantified using a fluorimeter.

(c) Various enzyme-substrate labels are available and U.S. Patent No. 4,275,149 provides a review of some of these. The enzyme generally catalyzes a chemical alteration of the chromogenic substrate which can be measured using various techniques. For example, the enzyme may catalyze a color change in a substrate, which can be measured spectrophotometrically. Alternatively, the enzyme may alter the fluorescence or chemiluminescence of the substrate. Techniques for quantifying a change in fluorescence are described above. The chemiluminescent substrate becomes electronically excited by a chemical reaction and may then emit light which can be measured (using a chemiluminometer, for example) or donates energy to a fluorescent acceptor. Examples of enzymatic labels include luciferases (e.g., firefly luciferase and bacterial luciferase; U.S. Patent No. 4,737,456), luciferin, 2,3-dihydrophthalazinediones, malate dehydrogenase, urease, peroxidase such as horseradish peroxidase (HRPO), alkaline phosphatase, beta-galactosidase, glucoamylase, lysozyme, saccharide oxidases (e.g., glucose oxidase, galactose oxidase, and glucose-6-phosphate dehydrogenase), heterocyclic oxidases (such as uricase and xanthine oxidase), lactoperoxidase, microperoxidase, and the like. Techniques for conjugating enzymes to antibodies are described in O'Sullivan et al., *Methods for the Preparation of Enzyme-Antibody Conjugates for use in Enzyme Immunoassay*, in *Methods in Enzym.* (ed J. Langone & H. Van Vunakis), Academic press, New York, 73:147-166 (1981).

Examples of enzyme-substrate combinations include, for example:

(i) Horseradish peroxidase (HRPO) with hydrogen peroxide as a substrate, wherein the hydrogen peroxidase oxidizes a dye precursor (e.g., orthophenylene diamine (OPD) or 3,3',5,5'-tetramethyl benzidine hydrochloride (TMB));

(ii) alkaline phosphatase (AP) with para-Nitrophenyl phosphate as chromogenic substrate; and

(iii) beta-D-galactosidase (beta-D-Gal) with a chromogenic substrate (e.g., p-nitrophenyl-beta-D-galactosidase) or fluorogenic substrate 4-

methyumbelliferyl-beta-D-galactosidase.

Numerous other enzyme-substrate combinations are available to those skilled in the art. For a general review of these, see U.S. Patent Nos. 4,275,149 and 4,318,980.

Sometimes, the label is indirectly conjugated with the antibody variant. The skilled artisan will be aware of various techniques for achieving this. For example, the antibody variant can be conjugated with biotin and any of the three broad categories of labels mentioned above can be conjugated with avidin, or vice versa. Biotin binds selectively to avidin and thus, the label can be conjugated with the antibody variant in this indirect manner. Alternatively, to achieve indirect conjugation of the label with the antibody variant, the antibody variant is conjugated with a small hapten (e.g., digoxin) and one of the different types of labels mentioned above is conjugated with an anti-hapten antibody variant (e.g., anti-digoxin antibody). Thus, indirect conjugation of the label with the antibody variant can be achieved.

In another embodiment of the invention, the antibody variant need not be labeled, and the presence thereof can be detected using a labeled antibody which binds to the antibody variant.

The antibodies of the present invention may be employed in any known assay method, such as competitive binding assays, direct and indirect sandwich assays, and immunoprecipitation assays. Zola, *Monoclonal Antibodies: A Manual of Techniques*, pp.147-158 (CRC Press, Inc. 1987).

Competitive binding assays rely on the ability of a labeled standard to compete with the test sample analyze for binding with a limited amount of antibody variant. The amount of antigen in the test sample is inversely proportional to the amount of standard that becomes bound to the antibodies. To facilitate determining the amount of standard that becomes bound, the antibodies generally are insolubilized before or after the competition, so that the standard and analyze that are bound to the antibodies may conveniently be separated from the standard and analyze which remain unbound.

Sandwich assays involve the use of two antibodies, each capable of binding to a different immunogenic portion, or epitope, of the protein to be detected. In a sandwich assay, the test sample analyze is bound by a first antibody which is immobilized on a solid support, and thereafter a second antibody binds to the analyze, thus forming an insoluble three-part

5 complex. See, e.g., US Pat No. 4,376,110. The second antibody may itself
be labeled with a detectable moiety (direct sandwich assays) or may be
10 measured using an anti-immunoglobulin antibody that is labeled with a
detectable moiety (indirect sandwich assay). For example, one type of
5 sandwich assay is an ELISA assay, in which case the detectable moiety is an
enzyme.

15 For immunohistochemistry, the tumor sample may be fresh or frozen or
may be embedded in paraffin and fixed with a preservative such as formalin,
for example.

20 The antibodies may also be used for in vivo diagnostic assays.
Generally, the antibody variant is labeled with a radionuclide (such as
111In, 99Tc, 14C, 131I, 125I, 3H, 32P or 35S) so that the tumor can be
localized using immunoscintigraphy.

E. Diagnostic Kits

25 As a matter of convenience, the antibody variant of the present
invention can be provided in a kit, i.e., a packaged combination of
reagents in predetermined amounts with instructions for performing the
diagnostic assay. Where the antibody variant is labeled with an enzyme, the
kit will include substrates and cofactors required by the enzyme (e.g., a
30 substrate precursor which provides the detectable chromophore or
fluorophore). In addition, other additives may be included such as
stabilizers, buffers (e.g., a block buffer or lysis buffer) and the like.
The relative amounts of the various reagents may be varied widely to
35 provide for concentrations in solution of the reagents which substantially
optimize the sensitivity of the assay. Particularly, the reagents may be
provided as dry powders, usually lyophilized, including excipients which on
dissolution will provide a reagent solution having the appropriate
concentration.

F. In Vivo Uses for the Antibody Variant

40 For therapeutic applications, the antibody variants of the invention
are administered to a mammal, preferably a human, in a pharmaceutically
acceptable dosage form such as those discussed above, including those that
45 may be administered to a human intravenously as a bolus or by continuous
infusion over a period of time, by intramuscular, intraperitoneal, intra-
cerebrospinal, subcutaneous, intra-articular, intrasynovial, intrathecal,
35 oral, topical, or inhalation routes. The antibodies also are suitably
administered by intra-tumoral, peri-tumoral, intra-lesional, or peri-

5

lesional routes, to exert local as well as systemic therapeutic effects. The intra-peritoneal route is expected to be particularly useful, for example, in the treatment of ovarian tumors. In addition, the antibody variant is suitably administered by pulse infusion, particularly with declining doses of the antibody variant. Preferably the dosing is given by injections, most preferably intravenous or subcutaneous injections, depending in part on whether the administration is brief or chronic.

10

5

15

10
15
20
25
30
35
40
45
50
55
60
65
70
75
80
85
90
95
100
105
110
115
120
125
130
135
140
145
150
155
160
165
170
175
180
185
190
195
200
205
210
215
220
225
230
235
240
245
250
255
260
265
270
275
280
285
290
295
300
305
310
315
320
325
330
335
340
345
350
355
360
365
370
375
380
385
390
395
400
405
410
415
420
425
430
435
440
445
450
455
460
465
470
475
480
485
490
495
500
505
510
515
520
525
530
535
540
545
550
555
560
565
570
575
580
585
590
595
600
605
610
615
620
625
630
635
640
645
650
655
660
665
670
675
680
685
690
695
700
705
710
715
720
725
730
735
740
745
750
755
760
765
770
775
780
785
790
795
800
805
810
815
820
825
830
835
840
845
850
855
860
865
870
875
880
885
890
895
900
905
910
915
920
925
930
935
940
945
950
955
960
965
970
975
980
985
990
995
1000
1005
1010
1015
1020
1025
1030
1035
1040
1045
1050
1055
1060
1065
1070
1075
1080
1085
1090
1095
1100
1105
1110
1115
1120
1125
1130
1135
1140
1145
1150
1155
1160
1165
1170
1175
1180
1185
1190
1195
1200
1205
1210
1215
1220
1225
1230
1235
1240
1245
1250
1255
1260
1265
1270
1275
1280
1285
1290
1295
1300
1305
1310
1315
1320
1325
1330
1335
1340
1345
1350
1355
1360
1365
1370
1375
1380
1385
1390
1395
1400
1405
1410
1415
1420
1425
1430
1435
1440
1445
1450
1455
1460
1465
1470
1475
1480
1485
1490
1495
1500
1505
1510
1515
1520
1525
1530
1535
1540
1545
1550
1555
1560
1565
1570
1575
1580
1585
1590
1595
1600
1605
1610
1615
1620
1625
1630
1635
1640
1645
1650
1655
1660
1665
1670
1675
1680
1685
1690
1695
1700
1705
1710
1715
1720
1725
1730
1735
1740
1745
1750
1755
1760
1765
1770
1775
1780
1785
1790
1795
1800
1805
1810
1815
1820
1825
1830
1835
1840
1845
1850
1855
1860
1865
1870
1875
1880
1885
1890
1895
1900
1905
1910
1915
1920
1925
1930
1935
1940
1945
1950
1955
1960
1965
1970
1975
1980
1985
1990
1995
2000
2005
2010
2015
2020
2025
2030
2035
2040
2045
2050
2055
2060
2065
2070
2075
2080
2085
2090
2095
2100
2105
2110
2115
2120
2125
2130
2135
2140
2145
2150
2155
2160
2165
2170
2175
2180
2185
2190
2195
2200
2205
2210
2215
2220
2225
2230
2235
2240
2245
2250
2255
2260
2265
2270
2275
2280
2285
2290
2295
2300
2305
2310
2315
2320
2325
2330
2335
2340
2345
2350
2355
2360
2365
2370
2375
2380
2385
2390
2395
2400
2405
2410
2415
2420
2425
2430
2435
2440
2445
2450
2455
2460
2465
2470
2475
2480
2485
2490
2495
2500
2505
2510
2515
2520
2525
2530
2535
2540
2545
2550
2555
2560
2565
2570
2575
2580
2585
2590
2595
2600
2605
2610
2615
2620
2625
2630
2635
2640
2645
2650
2655
2660
2665
2670
2675
2680
2685
2690
2695
2700
2705
2710
2715
2720
2725
2730
2735
2740
2745
2750
2755
2760
2765
2770
2775
2780
2785
2790
2795
2800
2805
2810
2815
2820
2825
2830
2835
2840
2845
2850
2855
2860
2865
2870
2875
2880
2885
2890
2895
2900
2905
2910
2915
2920
2925
2930
2935
2940
2945
2950
2955
2960
2965
2970
2975
2980
2985
2990
2995
3000
3005
3010
3015
3020
3025
3030
3035
3040
3045
3050
3055
3060
3065
3070
3075
3080
3085
3090
3095
3100
3105
3110
3115
3120
3125
3130
3135
3140
3145
3150
3155
3160
3165
3170
3175
3180
3185
3190
3195
3200
3205
3210
3215
3220
3225
3230
3235
3240
3245
3250
3255
3260
3265
3270
3275
3280
3285
3290
3295
3300
3305
3310
3315
3320
3325
3330
3335
3340
3345
3350
3355
3360
3365
3370
3375
3380
3385
3390
3395
3400
3405
3410
3415
3420
3425
3430
3435
3440
3445
3450
3455
3460
3465
3470
3475
3480
3485
3490
3495
3500
3505
3510
3515
3520
3525
3530
3535
3540
3545
3550
3555
3560
3565
3570
3575
3580
3585
3590
3595
3600
3605
3610
3615
3620
3625
3630
3635
3640
3645
3650
3655
3660
3665
3670
3675
3680
3685
3690
3695
3700
3705
3710
3715
3720
3725
3730
3735
3740
3745
3750
3755
3760
3765
3770
3775
3780
3785
3790
3795
3800
3805
3810
3815
3820
3825
3830
3835
3840
3845
3850
3855
3860
3865
3870
3875
3880
3885
3890
3895
3900
3905
3910
3915
3920
3925
3930
3935
3940
3945
3950
3955
3960
3965
3970
3975
3980
3985
3990
3995
4000
4005
4010
4015
4020
4025
4030
4035
4040
4045
4050
4055
4060
4065
4070
4075
4080
4085
4090
4095
4100
4105
4110
4115
4120
4125
4130
4135
4140
4145
4150
4155
4160
4165
4170
4175
4180
4185
4190
4195
4200
4205
4210
4215
4220
4225
4230
4235
4240
4245
4250
4255
4260
4265
4270
4275
4280
4285
4290
4295
4300
4305
4310
4315
4320
4325
4330
4335
4340
4345
4350
4355
4360
4365
4370
4375
4380
4385
4390
4395
4400
4405
4410
4415
4420
4425
4430
4435
4440
4445
4450
4455
4460
4465
4470
4475
4480
4485
4490
4495
4500
4505
4510
4515
4520
4525
4530
4535
4540
4545
4550
4555
4560
4565
4570
4575
4580
4585
4590
4595
4600
4605
4610
4615
4620
4625
4630
4635
4640
4645
4650
4655
4660
4665
4670
4675
4680
4685
4690
4695
4700
4705
4710
4715
4720
4725
4730
4735
4740
4745
4750
4755
4760
4765
4770
4775
4780
4785
4790
4795
4800
4805
4810
4815
4820
4825
4830
4835
4840
4845
4850
4855
4860
4865
4870
4875
4880
4885
4890
4895
4900
4905
4910
4915
4920
4925
4930
4935
4940
4945
4950
4955
4960
4965
4970
4975
4980
4985
4990
4995
5000
5005
5010
5015
5020
5025
5030
5035
5040
5045
5050
5055
5060
5065
5070
5075
5080
5085
5090
5095
5100
5105
5110
5115
5120
5125
5130
5135
5140
5145
5150
5155
5160
5165
5170
5175
5180
5185
5190
5195
5200
5205
5210
5215
5220
5225
5230
5235
5240
5245
5250
5255
5260
5265
5270
5275
5280
5285
5290
5295
5300
5305
5310
5315
5320
5325
5330
5335
5340
5345
5350
5355
5360
5365
5370
5375
5380
5385
5390
5395
5400
5405
5410
5415
5420
5425
5430
5435
5440
5445
5450
5455
5460
5465
5470
5475
5480
5485
5490
5495
5500
5505
5510
5515
5520
5525
5530
5535
5540
5545
5550
5555
5560
5565
5570
5575
5580
5585
5590
5595
5600
5605
5610
5615
5620
5625
5630
5635
5640
5645
5650
5655
5660
5665
5670
5675
5680
5685
5690
5695
5700
5705
5710
5715
5720
5725
5730
5735
5740
5745
5750
5755
5760
5765
5770
5775
5780
5785
5790
5795
5800
5805
5810
5815
5820
5825
5830
5835
5840
5845
5850
5855
5860
5865
5870
5875
5880
5885
5890
5895
5900
5905
5910
5915
5920
5925
5930
5935
5940
5945
5950
5955
5960
5965
5970
5975
5980
5985
5990
5995
6000
6005
6010
6015
6020
6025
6030
6035
6040
6045
6050
6055
6060
6065
6070
6075
6080
6085
6090
6095
6100
6105
6110
6115
6120
6125
6130
6135
6140
6145
6150
6155
6160
6165
6170
6175
6180
6185
6190
6195
6200
6205
6210
6215
6220
6225
6230
6235
6240
6245
6250
6255
6260
6265
6270
6275
6280
6285
6290
6295
6300
6305
6310
6315
6320
6325
6330
6335
6340
6345
6350
6355
6360
6365
6370
6375
6380
6385
6390
6395
6400
6405
6410
6415
6420
6425
6430
6435
6440
6445
6450
6455
6460
6465
6470
6475
6480
6485
6490
6495
6500
6505
6510
6515
6520
6525
6530
6535
6540
6545
6550
6555
6560
6565
6570
6575
6580
6585
6590
6595
6600
6605
6610
6615
6620
6625
6630
6635
6640
6645
6650
6655
6660
6665
6670
6675
6680
6685
6690
6695
6700
6705
6710
6715
6720
6725
6730
6735
6740
6745
6750
6755
6760
6765
6770
6775
6780
6785
6790
6795
6800
6805
6810
6815
6820
6825
6830
6835
6840
6845
6850
6855
6860
6865
6870
6875
6880
6885
6890
6895
6900
6905
6910
6915
6920
6925
6930
6935
6940
6945
6950
6955
6960
6965
6970
6975
6980
6985
6990
6995
7000
7005
7010
7015
7020
7025
7030
7035
7040
7045
7050
7055
7060
7065
7070
7075
7080
7085
7090
7095
7100
7105
7110
7115
7120
7125
7130
7135
7140
7145
7150
7155
7160
7165
7170
7175
7180
7185
7190
7195
7200
7205
7210
7215
7220
7225
7230
7235
7240
7245
7250
7255
7260
7265
7270
7275
7280
7285
7290
7295
7300
7305
7310
7315
7320
7325
7330
7335
7340
7345
7350
7355
7360
7365
7370
7375
7380
7385
7390
7395
7400
7405
7410
7415
7420
7425
7430
7435
7440
7445
7450
7455
7460
7465
7470
7475
7480
7485
7490
7495
7500
7505
7510
7515
7520
7525
7530
7535
7540
7545
7550
7555
7560
7565
7570
7575
7580
7585
7590
7595
7600
7605
7610
7615
7620
7625
7630
7635
7640
7645
7650
7655
7660
7665
7670
7675
7680
7685
7690
7695
7700
7705
7710
7715
7720
7725
7730
7735
7740
7745
7750
7755
7760
7765
7770
7775
7780
7785
7790
7795
7800
7805
7810
7815
7820
7825
7830
7835
7840
7845
7850
7855
7860
7865
7870
7875
7880
7885
7890
7895
7900
7905
7910
7915
7920
7925
7930
7935
7940
7945
7950
7955
7960
7965
7970
7975
7980
7985
7990
7995
8000
8005
8010
8015
8020
8025
8030
8035
8040
8045
8050
8055
8060
8065
8070
8075
8080
8085
8090
8095
8100
8105
8110
8115
8120
8125
8130
8135
8140
8145
8150
8155
8160
8165
8170
8175
8180
8185
8190
8195
8200
8205
8210
8215
8220
8225
8230
8235
8240
8245
8250
8255
8260
8265
8270
8275
8280
8285
8290
8295
8300
8305
8310
8315
8320
8325
8330
8335
8340
8345
8350
8355
8360
8365
8370
8375
8380
8385
8390
8395
8400
8405
8410
8415
8420
8425
8430
8435
8440
8445
8450
8455
8460
8465
8470
8475
8480
8485
8490
8495
8500
8505
8510
8515
8520
8525
8530
8535
8540
8545
8550
8555
8560
8565
8570
8575
8580
8585
8590
8595
8600
8605
8610
8615
8620
8625
8630
8635
8640
8645
8650
8655
8660
8665
8670
8675
8680
8685
8690
8695
8700
8705
8710
8715
8720
8725
8730
8735
8740
8745
8750
8755
8760
8765
8770
8775
8780
8785
8790
8795
8800
8805
8810
8815
8820
8825
8830
8835
8840
8845
8850
8855
8860
8865
8870
8875
8880
8885
8890
8895
8900
8905
8910
8915
8920
8925
8930
8935
8940
8945
8950
8955
8960
8965
8970
8975
8980
8985
8990
8995
9000
9005
9010
9015
9020
9025
9030
9035
9040
9045
9050
9055
9060
9065
9070
9075
9080
9085
9090
9095
9100
9105
9110
9115
9120
9125
9130
9135
9140
9145
9150
9155
9160
9165
9170
9175
9180
9185
9190
9195
9200
9205
9210
9215
9220
9225
9230
9235
9240
9245
9250
9255
9260
9265
9270
9275
9280
9285
9290
9295
9300
9305
9310
9315
9320
9325
9330
9335
9340
9345
9350
9355
9360
9365
9370
9375
9380
9385
9390
9395
9400
9405
9410
9415
9420
9425
9430
9435
9440
9445
9450
9455
9460
9465
9470
9475
9480
9485
9490
9495
9500
9505
9510
9515
9520
9525
9530
9535
9540
9545
9550
9555
9560
9565
9570
9575
9580
9585
9590
9595
9600
9605
9610
9615
9620
9625
9630
9635
9640
9645
9650
9655
9660
9665
9670
9675
9680
9685
9690
9695
9700
9705
9710
9715
9720
9725
9730
9735
9740
9745
9750
9755
9760
9765
9770
9775
9780
9785
9790
9795
9800
9805
9810
9815
9820
9825
9830
9835
9840
9845
9850
9855
9860
9865
9870
9875
9880
9885
9890
9895
9900
9905
9910
9915
9920
9925
9930
9935
9940
9945
9950
9955
9960
9965
9970
9975
9980
9985
9990
9995
10000
10005
10010
10015
10020
10025
10030
10035
10040
10045
10050
10055
10060
10065
10070
10075
10080
10085
10090
10095
10100
10105
10110
10115
10120
10125
10130
10135
10140
10145
10150
10155
10160
10165
10170
10175
10180

nephrotic syndrome, preeclampsia, ascites, pericardial effusion (such as that associated with pericarditis), and pleural effusion.

Age-related macular degeneration (AMD) is a leading cause of severe visual loss in the elderly population. The exudative form of AMD is characterized by choroidal neovascularization and retinal pigment epithelial cell detachment. Because choroidal neovascularization is associated with a dramatic worsening in prognosis, the VEGF antibodies of the present invention are expected to be especially useful in reducing the severity of AMD.

Depending on the type and severity of the disease, about 1 µg/kg to 15 mg/kg (e.g., 0.1-20mg/kg) of antibody variant is an initial candidate dosage for administration to the patient, whether, for example, by one or more separate administrations, or by continuous infusion. A typical daily dosage might range from about 1 µg/kg to 100 mg/kg or more, depending on the factors mentioned above. For repeated administrations over several days or longer, depending on the condition, the treatment is sustained until a desired suppression of disease symptoms occurs. However, other dosage regimens may be useful. The progress of this therapy is easily monitored by conventional techniques and assays.

The antibody variant composition will be formulated, dosed, and administered in a fashion consistent with good medical practice. Factors for consideration in this context include the particular disorder being treated, the particular mammal being treated, the clinical condition of the individual patient, the cause of the disorder, the site of delivery of the agent, the method of administration, the scheduling of administration, and other factors known to medical practitioners. The "therapeutically effective amount" of the antibody variant to be administered will be governed by such considerations, and is the minimum amount necessary to prevent, ameliorate, or treat a disease or disorder. The antibody variant need not be, but is optionally formulated with one or more agents currently used to prevent or treat the disorder in question. The effective amount of such other agents depends on the amount of antibody variant present in the formulation, the type of disorder or treatment, and other factors discussed above. These are generally used in the same dosages and with administration routes as used hereinbefore or about from 1 to 99% of the heretofore employed dosages.

G. Articles of Manufacture

5

10

15

20

25

In another embodiment of the invention, an article of manufacture containing materials useful for the treatment of the disorders described above is provided. The article of manufacture comprises a container and a label. Suitable containers include, for example, bottles, vials, syringes, and test tubes. The containers may be formed from a variety of materials such as glass or plastic. The container holds a composition which is effective for treating the condition and may have a sterile access port (for example the container may be an intravenous solution bag or a vial having a stopper pierceable by a hypodermic injection needle). The active agent in the composition is the antibody variant. The label on, or associated with, the container indicates that the composition is used for treating the condition of choice. The article of manufacture may further comprise a second container comprising a pharmaceutically-acceptable buffer, such as phosphate-buffered saline, Ringer's solution and dextrose solution. It may further include other materials desirable from a commercial and user standpoint, including other buffers, diluents, filters, needles, syringes, and package inserts with instructions for use.

Example 1

30

35

40

In this example, antibody variants containing randomized peptide inserts within the antibody CDRs are prepared by phage display which substantially improve the affinity of a humanized Fab for VEGF. Crystallography suggests that these changes result in an increased contact area with antigen.

VEGF:Fab X-ray Co-Crystal Structure: A crystal structure of the complex between the VEGF antigen and anti-VEGF parent antibody was prepared as described in Muller et al., *Structure* 6(9):1153-1167 (1998). The conclusion that the three VH CDRs are the main determinants of Fab binding to VEGF is supported by the high-resolution crystal structure of the VEGF:Fab (v36) complex. In addition, the major energetic determinants largely coincide with the principal contacting residues of the Fab in the complex.

45

Several randomized libraries were designed with a peptide insertion placed in the antigen-contacting CDRs which, from the crystal structure, were expected to increase the potential contact between the antibody and the antigen.

50

Design of CDR Random Loop-Insertion Libraries: Based upon inspection

55

of the VEGF:Fab crystal structure, it was postulated that additional contacts, contributing additional binding energy between the Fab and VEGF, could be generated through the addition of peptide inserts within one or more CDRs of the Fab. Because the nature and relative contributions of such additional interactions would be difficult to predict, randomized loop sequences (Xn) were directly inserted into each of the four CDRs proximal to the existing VEGF binding site using NNS codons, and a frameshifted Fab vector as template. The length of loop was chosen based upon distances in the crystal structure between exit/entry points of the loop on the hypervariable region and possible interaction sites on the surface of VEGF. In addition, one or more residues within each loop were deleted in some of these templates, as judged necessary to accommodate the new peptide loop.

Three such loops were designed for VH1, including insertions of 4, 5, or 6 residues between Y27 and T28. In VH2, two inserted peptides of 3 or 4 residues were placed between Y54 and T55. Also in VH2, a 6-residue random peptide was used to replace residues T55 and H56. In VH3, a 4-residue or 5-residue peptide was used to replace G104, and a 5-residue or 6-residue peptide was used to replace residues G104 and S105. Finally, in VL3, a random peptide of either 4 or 6 residues was inserted between S92 and T93.

Second-Generation Selections of anti-VEGF Libraries: Templates for random mutagenesis were constructed starting from the Fab-g3 phagemid pY0192 (WO98/45331) and frameshift oligonucleotides (which prevent expression of a functional template Fab): YC-82, YC-85, YC-89, YC-92, YC-94, and YC-97 (Table 1).

Table 1

Frameshift oligos for CDR-insert template mutagenesis

Oligo #	Region	Sequence	SEQ. ID NO:
YC-82	VL3	C TGT CAA CAG TAT AGC T ACC GTG CCG TGG ACG	SEQ.ID NO:1
YC-85	VH1	GCA GCT TCT GGC TAT G ACC TTC ACC AAC TAT G	SEQ.ID NO:2
YC-89	VH2	GA TGG ATT AAC ACC TAT G ACC GGT GAA CCG ACC	SEQ.ID NO:3
YC-92	VH2	GA TGG ATT AAC ACC TAT T GAA CCG ACC TAT GCT G	SEQ.ID NO:4
YC-94	VH3	G TAC CCG CAC TAT TAT G AGC AGC CAC TGG TAT TTC	SEQ.ID NO:5
YC-97	VH3	G TAC CCG CAC TAT TAT G AGC CAC TGG TAT TTC	SEQ.ID NO:6

The corresponding randomization oligonucleotides (which employ NNS at the sites targeted for randomization) were YC-83, YC-84 in VL3; YC-86, YC-

87, YC-88 in VH1; YC-90, YC91 and YC-93 in VH2; and YC-95, YC-96, YC-98, YC-99 in VH3. See Table 2 below.

Table 2

Random oligos for CDR-insert library constructions

Oligo #	Region	(Comments)	Sequence	SEQ ID NO:
YC-83	VL3	(insert 4 residues)	C TGT CAA CAG TAT AGC NNS NNS NNS NNS ACC GTG CCG TGG ACG	SEQ. ID NO:7
YC-84	VL3	(insert 6 residues)	C TGT CAA CAG TAT AGC NNS NNS NNS NNS NNS NNS ACC GTG CCG TGG ACG	SEQ. ID NO:8
YC-86	VH1	(insert 4 residues)	GCA GCT TCT GGC TAT NNS NNS NNS NNS ACC TTC ACC AAC TAT G	SEQ. ID NO:9
YC-87	VH1	(insert 5 residues)	GCA GCT TCT GGC TAT NNS NNS NNS NNS NNS ACC TTC ACC AAC TAT G	SEQ. ID NO:10
YC-88	VH1	(insert 6 residues)	GCA GCT TCT GGC TAT NNS NNS NNS NNS NNS NNS ACC TTC ACC AAC TAT G	SEQ. ID NO:11
YC-90	VH2	(insert 3 residues)	GA TGG ATT AAC ACC TAT NNS NNS NNS ACC GGT GAA CCG ACC	SEQ. ID NO:12
YC-91	VH2	(insert 4 residues)	GA TGG ATT AAC ACC TAT NNS NNS NNS NNS ACC GGT GAA CCG ACC	SEQ. ID NO:13
YC-93	VH2	(insert 6 residues)	GA TGG ATT AAC ACC TAT NNS NNS NNS NNS NNS NNS GAA CCG ACC TAT GCT G	SEQ. ID NO:14
YC-95	VH3	(insert 4 residues)	G TAC CCG CAC TAT TAT NNS NNS NNS NNS AGC AGC CAC TGG TAT TTC	SEQ. ID NO:15
YC-96	VH3	(insert 5 residues)	G TAC CCG CAC TAT TAT NNS NNS NNS NNS NNS AGC AGC CAC TGG TAT TTC	SEQ. ID NO:16
YC-98	VH3	(insert 5 residues)	G TAC CCG CAC TAT TAT NNS NNS NNS NNS NNS AGC CAC TGG TAT TTC	SEQ. ID NO:17
YC-99	VH3	(insert 6 residues)	G TAC CCG CAC TAT TAT NNS NNS NNS NNS NNS NNS AGC CAC TGG TAT TTC	SEQ. ID NO:18

The resulting transformants yielded libraries with complexities ranging from 6×10^7 to 5×10^8 suggesting that the libraries were comprehensive in covering all possible variants.

Each library was sorted separately for the first round; thereafter, libraries with the same site of insertion were combined and sorted together as one. Therefore, library YC-83 was combined with library YC-84; library YC-86 with libraries YC-87 and YC-88; library YC-90 with YC-91; library YC-95 with YC-96; and library YC-98 with YC-99. These libraries were sorted essentially as described in WO98/45331, except the incubation with PBS/TWEEN 20® buffer after phage binding was carried out as described in Table 3.

5

Table 3
Conditions for secondary selections of Fab variants

10

15

round of selection	incubation time (hr)	incubation solution	incubation temp. (°C)
1	0	0	room temp.
2	1	ELISA buffer	room temp.
3	2	1 μ M VEGF/ELISA	room temp.
4	18	1 μ M VEGF/ELISA	room temp.
5	37	1 μ M VEGF /ELISA	room temp.
6	17hr@R.T./30h@37°C	same as above	room temp./37°C
7	63	same as above	37°C
8	121	same as above	37°C

20

5 ELISA buffer contained 0.5% bovine serum albumin and 0.05% TWEEN 20® in PBS. VEGF was included in the incubation buffer to minimize rebinding of phage to VEGF coated on the surface of the plate.

25

Sorting of some of these libraries yielded VEGF-binding phage enrichments over 5 to 8 rounds of selection. After five to eight rounds of selections, ten to twenty clones from each library were isolated from carbenicillin containing plates harboring *E. coli* (XL1) colonies which had been infected with an eluted phage pool. Colonies were isolated and grown with helper phage to obtain single-stranded DNA for sequencing. Clones were picked from those libraries that enriched for DNA sequencing. The results are shown in Table 4. Libraries showing no enrichment were not sequenced.

30

35

Table 4
Summary of CDR Insertion Libraries

40

45

Oligos		CDR	Site of Insertion	No. of added residues	
Stop oligo	Insert oligo			Net	Total
YC-85	YC-86	H1	Y27^T28	4	4
YC-85	YC-87	H1	Y27^T28	5	5
YC-85	YC-88	H1	Y27^T28	6	6
YC-89	YC-90	H2	Y54^T55	3	3
YC-89	YC-91	H2	Y54^T55	4	4
YC-92	YC-93	H2	Y54^E57	4	6
YC-94	YC-95	H3	Y103^S105	3	4
YC-94	YC-96	H3	Y103^S105	4	5
YC-97	YC-98	H3	Y103^S106	3	5
YC-97	YC-99	H3	Y103^S106	4	6
YC-82	YC-83	L3	S92^T93	4	4
YC-82	YC-84	L3	S92^T93	6	6

50

20 For VH1, only library YC-86 showed enrichment. Sequencing revealed

55

that, although a 4-residue insert was designed in this library, all of the sequenced clones contained no net insertion, but instead point mutations at T28 and F29. This suggests that this antibody is relatively intolerant of insertions in this hypervariable region.

A similar result was seen for the VH2 libraries, where only library YC-90 showed enrichment. Again, clones found were either wild-type (Y0192) or a point mutant, Y54W. This suggests that this antibody is also relatively intolerant of insertions in the VH2 CDR.

Again, a similar result was obtained for the VL3 libraries. In this case, only library YC-83 showed enrichment, and the selected clones had point mutations at T93 and/or V94, rather than the designed insertion. This suggests that this antibody is also relatively intolerant of insertions in the VL3 CDR.

In contrast, two VH3 libraries showed enrichment: YC-95 and YC-98. Moreover, sequencing of selected clones showed that the Fab variants indeed contained insertion sequences.

Amino acid sequences of anti-VEGF variants from the various libraries are shown in Tables 5-15 below. The sequence of the randomized region only is shown as deduced from DNA sequencing. Sites where randomized inserted sequences were made are shown in bold. An asterisk denotes a contaminating phagemid from another library.

Table 5

Protein sequences of anti-VEGF variants from library YC-86
Round 7 (VEGF eluted phage)

Name	VH1 sequence (residues 26-35)	SEQ ID NO:	(# clones/10)
Y0241-1	GYDPTNYGIN	SEQ. ID NO:19	4
Y0241-6	GYDYTNNGIN	SEQ. ID NO:20	3
Y0241-7	GYDWTNYGIN	SEQ. ID NO:21	3

Table 6

Protein sequences of anti-VEGF variants from library YC-90
Round 7 (VEGF eluted phage)

Name	VH2 sequence (residues 50-62)	SEQ ID NO:	(# clones/10)
YC242-1	WINTWTGEPTYAA	SEQ. ID NO:22	4
*Y0192			6

Table 7

Protein sequences of anti-VEGF variants from library YC-83
Round 7 (VEGF eluted phage)

Name	VL3 sequence (residues 89-97)	SEQ ID NO:	(# clones/9)
Y0241-2	QQYSATPWT	SEQ. ID NO:23	1
Y0241-3	QQYSNPWT	SEQ. ID NO:24	3
Y0241-4	QQYSAVPWT	SEQ. ID NO:25	4
Y0241-5	QQYSVPWT	SEQ. ID NO:26	1

Table 8

Protein sequences of anti-VEGF variants from library YC-95
Round 5 (VEGF eluted phage)

Name	VH3 sequence (residues 99-111)+ insertions	SEQ ID NO:	(# clones/10)
Y0228-1	YPHYVAKERSHHWYFDV	SEQ. ID NO:27	1
Y0228-2	YPHYVGETSSHWYFDV	SEQ. ID NO:28	1
Y0228-3	YPHYVARDRSSHWYFDV	SEQ. ID NO:29	1
Y0228-4	YPHYVERDGKSSHWYFDV	SEQ. ID NO:30	1
Y0228-5	YPHYVRNEKSSHWYFDV	SEQ. ID NO:31	1
Y0228-6	YPHYVVEQSSHWYFDV	SEQ. ID NO:32	1
Y0228-7	YPHYVQRDRSSHWYFDV	SEQ. ID NO:33	1
Y0228-8	YPHYVQKQKSSHWYFDV	SEQ. ID NO:34	1
Y0228-9	YPHYVQNEGPSSHWYFDV	SEQ. ID NO:35	1
Y0228-10	YPHYVGNRRSSHWYFDV	SEQ. ID NO:36	1

Table 9

Protein sequences of anti-VEGF variants from library YC-95
Round 5 (HCl eluted phage)

Name	VH3 sequence (residues 99-111)+ insertions	SEQ ID NO:	(# clones/10)
Y0229-1	YPHYVTEKSSHWYFDV	SEQ. ID NO:37	1
Y0229-2	YPHYVLKDRSSHWYFDV	SEQ. ID NO:38	1
Y0229-4	YPHYVQDEKSSHWYFDV	SEQ. ID NO:39	1
Y0229-5	YPHYVVEKSSHWYFDV	SEQ. ID NO:40	1
Y0229-6	YPHYVRDEKSSHWYFDV	SEQ. ID NO:41	1
Y0229-7	YPHYVTDKSSHWYFDV	SEQ. ID NO:42	1
Y0229-8	YPHYVTRGGSSHWYFDV	SEQ. ID NO:43	1
Y0229-9	YPHYVLNDRSSHWYFDV	SEQ. ID NO:44	1
Y0229-10	YPHYVTRDRSSHWYFDV	SEQ. ID NO:45	1
*Y0239-1			1

Table 10

Protein sequences of anti-VEGF variants from library YC-95

Round 7 (HCl eluted phage)

Name	VH3 sequence (residues 99-111)+ insertions	SEQ ID NO:	(# clones/10)
Y0239-1	YPHYVRNERSHWFYFDV	SEQ. ID NO:46	1
Y0239-2	YPHYVKNDKSSHWFYFDV	SEQ. ID NO:47	1
Y0239-3	YPHYVLADRSSHWFYFDV	SEQ. ID NO:48	1
Y0239-4	YPHYVNERSSHWFYFDV	SEQ. ID NO:49	1
Y0239-5	YPHYVLKDKSSHWFYFDV	SEQ. ID NO:50	1
Y0239-6	YPHYVLKGRSSHWFYFDV	SEQ. ID NO:51	1
Y0239-7	YPHYVERDGRSSHWFYFDV	SEQ. ID NO:52	1
Y0239-8	YPHYVLKGRSSHWFYFDV	SEQ. ID NO:53	1
Y0239-9	YPHYVLGESSHWFYFDV	SEQ. ID NO:54	1
Y0239-10	YPHYVLGKSSHWFYFDV	SEQ. ID NO:55	1

Table 11

Protein sequences of anti-VEGF variants from library YC-95

Round 8 (HCl eluted phage)

Name	VH3 sequence (residues 99-111)+ insertions	SEQ ID NO:	(# clones/10)
Y0261-1	YPHYVLKDRSSHWFYFDV	SEQ. ID NO:56	2
Y0261-2	YPHYVLKDMSSHWFYFDV	SEQ. ID NO:57	2
*Y0239-4			1
*Y0239-9			5

Table 12

Protein sequences of anti-VEGF variants from library YC-98

Round 5 (VEGF eluted phage)

Name	VH3 sequence (residues 99-111)+ insertions	SEQ ID NO:	(# clones/10)
Y0228-11	YPHYVEKQRKSHWFYFDV	SEQ. ID NO:58	1
Y0228-12	YPHYVKEDEKSHWFYFDV	SEQ. ID NO:59	1
Y0228-13	YPHYVSHQKRSHWFYFDV	SEQ. ID NO:60	1
Y0228-14	YPHYVSGERESHWFYFDV	SEQ. ID NO:61	1
Y0228-15	YPHYVQSEGRSHWFYFDV	SEQ. ID NO:62	1
Y0228-16	YPHYVSVEGGSHWFYFDV	SEQ. ID NO:63	1
Y0228-17	YPHYVPSPRGSHWFYFDV	SEQ. ID NO:64	1
Y0228-18	YPHYVQRNGKSHWFYFDV	SEQ. ID NO:65	1
Y0228-19	YPHYVAREGGSHWFYFDV	SEQ. ID NO:66	1
Y0228-20	YPHYVSNERKSHWFYFDV	SEQ. ID NO:67	1

Table 13

Protein sequences of anti-VEGF variants from library YC-98
Round 5 (HCl eluted phage)

Name	VH3 sequence (residues 99-111)+ insertions	SEQ ID NO:	(# clones/10)
Y0229-11	YPHYRGRKSHWYFDV	SEQ. ID NO:68	1
Y0229-12	YPHYRDERKSHWYFDV	SEQ. ID NO:69	1
Y0229-13	YPHYRGRKSHWYFDV	SEQ. ID NO:70	1
Y0229-14	YPHYRGRKSHWYFDV	SEQ. ID NO:71	1
Y0229-15	YPHYRGRKSHWYFDV	SEQ. ID NO:72	1
Y0229-16	YPHYRGRKSHWYFDV	SEQ. ID NO:73	1
Y0229-17	YPHYRGRKSHWYFDV	SEQ. ID NO:74	1
Y0229-18	YPHYRGRKSHWYFDV	SEQ. ID NO:75	1
Y0229-19	YPHYRGRKSHWYFDV	SEQ. ID NO:76	1

Table 14

Protein sequences of anti-VEGF variants from library YC-98
Round 7 (HCl eluted phage)

Name	VH3 sequence (residues 99-111)+ insertions	SEQ ID NO:	(# clones/10)
Y0239-11	YPHYRDERKSHWYFDV	SEQ. ID NO:77	1
Y0239-12	YPHYRGRKSHWYFDV	SEQ. ID NO:78	1
Y0239-13	YPHYRGRKSHWYFDV	SEQ. ID NO:79	1
Y0239-14	YPHYRGRKSHWYFDV	SEQ. ID NO:80	1
Y0239-15	YPHYRGRKSHWYFDV	SEQ. ID NO:81	1
Y0239-16	YPHYRGRKSHWYFDV	SEQ. ID NO:82	1
Y0239-17	YPHYRGRKSHWYFDV	SEQ. ID NO:83	1
Y0239-18	YPHYRGRKSHWYFDV	SEQ. ID NO:84	1
Y0239-19	YPHYRGRKSHWYFDV	SEQ. ID NO:85	2
Y0240-1	YPHYRGRKSHWYFDV	SEQ. ID NO:86	1

Table 15

Protein sequences of anti-VEGF variants from library YC-98
Round 8 (HCl eluted phage)

Name	VH3 sequence (residues 99-111)+ insertions	SEQ ID NO:	(# clones/10)
Y0261-4	YPHYRGRKSHWYFDV	SEQ. ID NO:87	1
Y0261-5	YPHYRGRKSHWYFDV	SEQ. ID NO:88	1
Y0261-6	YPHYRGRKSHWYFDV	SEQ. ID NO:89	1
Y0261-7	YPHYRGRKSHWYFDV	SEQ. ID NO:90	1
Y0261-8	YPHYRGRKSHWYFDV	SEQ. ID NO:91	1
Y0239-19	YPHYRGRKSHWYFDV	SEQ. ID NO:92	1
*Y0239-13			1
*Y0239-16			3

In order to quantify relative antigen-binding affinities, several anti-VEGF variants' DNA were transformed into *E. coli* strain 3438,

expressed as Fab, and purified by passing the periplasmic shockate through a protein G column (Pharmacia) as described in WO98/45331.

CDR combination Variant Y0313-2: An attempt was made to improve antigen binding affinity by combining a previously discovered CDR VH2 mutation with an insertion variant described here. A mutagenic oligonucleotide, YC-107 (Table 16) was used to combine insertion mutations found in CDR VH3, from clone Y0239-19, with VH2 CDR mutations T28D/N31H from clone Y0243-1 (WO98/45331) of CDR VH2.

Table 16

Mutagenesis oligo for adding a CDR insertion peptide

Oligo #	Region	(Comments)	Sequence	SEQ. ID NO:
YC-107	VH3	(insert VNERK from library YC-98)	TAC CCG CAC TAT TAT GTC AAC GAG CGG AAG AGC CAC TGG TAT TTC	SEQ. ID NO:93

The resulting combined CDR variant was designated YC313-2. A Fab protein sample was prepared as described above for BIACORE™ analysis.

BIACORE™ Analysis: The VEGF-binding affinities of Fab fragments were calculated from association and dissociation rate constants measured using a BIACORE™-2000 surface plasmon resonance system (BIACORE™, Inc., Piscataway, NJ). A biosensor chip was activated for covalent coupling of VEGF using N-ethyl-N'-(3-dimethylaminopropyl)-carbodiimide hydrochloride (EDC) and N-hydroxysuccinimide (NHS) according to the supplier's (BIACORE™, Inc., Piscataway, NJ) instructions. VEGF(8-109) was buffered exchanged into 20 mM sodium acetate, pH 4.8 and diluted to approximately 50 µg/mL. Aliquots of VEGF were injected at a flow rate of 2 µL/min to achieve approximately 700-1400 response units (RU) of coupled protein. A solution of 1 M methanolamine was injected as a blocking agent.

For kinetics measurements, two-fold serial dilutions of Fab were injected in PBS/TWEEN buffer (0.05% TWEEN 20™ in phosphate buffered saline) at 25°C at a flow rate of 10 µL/min. Equilibrium dissociation constants, Kd's from SPR measurements were calculated as koff/kon (Table 17).

Table 17

Kinetics of Fab-VEGF binding from BIACORE™ measurements.

Variant	$K_{on}(10^4/M/s)$	$k_{off}(10^{-4}/s)$	$K_d(nM)$	$K_d(wt)/K_d(mut)$
Y0192	4.1	1.21	2.9	-1-
Y0241-4	4.4	1.41	3.2	0.9
Y0241-7	4.6	1.28	3.0	1.0
Y0241-6	4.7	1.29	2.7	1.1
Y0242-1	4.7	0.86	1.8	1.6
Y0239-19	3.6	0.10	0.30	9.7
Y0239-8	3.8	0.18	0.50	5.8
Y0240-1	2.5	0.13	0.50	5.8
Y0239-2	3.6	1.64	4.6	0.6
Y0239-12	5.7	0.34	0.6	4.8
Y0239-9	3.97	0.19	0.5	6.0
YC261-6	4.4	0.25	0.6	5.0
YC313-2	3.11	0.11	0.36	8.0

Results of SPR measurements demonstrated that affinity is mainly enhanced through a slower dissociation rate (as opposed to faster association).

For the insertion variant Y0239-19, an approximately 10-fold improvement in binding affinity was observed (Table 17). However, addition of the VH1 mutations did not further improve affinity, as indicated for the variant Y0313-2.

Cell-Based Assay of VEGF: Two Fab variants of the anti-VEGF antibody were tested for their ability to antagonize VEGF (recombinant; version 1-165) in induction of the growth of HuVECs (human umbilical vein endothelial cells). The alamar blue assay (H. Gazzano-Santoro, et al. *J Immunol Methods* 202:163-171 (1997)) was used to measure the metabolic activity of cells in response to VEGF.

Two Fab variants of the anti-VEGF antibody were tested for their ability to antagonize VEGF (recombinant; version 1-165) activity in induction of the growth of HuVECs (human umbilical vein endothelial cells). HuVEC cells are seeded (1500/well) in a 96 well microtiter plate in complete medium (Cell Systems, Kirkland, WA) that has been coated with Cell Systems attachment factor. The cells are allowed to attach for 24 hrs. On day 2, VEGF and Fab are diluted in assay medium (DMEM/F12 +penicillin/streptomycin, 0.1% gelatin). For the antibody experiments, a constant concentration of 5 ng/ml VEGF is added to all the wells followed by the addition of various concentrations of anti-VEGF Fab (approximately 10 µg/ml and dilutions). The VEGF and Fab incubate with the HUVEC cells for 2 days, after which 25 µl of alamar blue is added. Following a 4 hr

incubation period, fluorescence is read on a Cytofluor Fluorescence Plate reader. The media used for these assays is from Cell Systems.

The results (Figure 2) show that the insertion variant Y0313-2 Fab has roughly 100-fold enhanced potency over the original humanized antibody, F(ab)-12.

Crystallization and X-Ray Structure Determination of the Insert-Fab Y0313-2 in complex with VEGF: Crystals of VEGF in complex with the Fab fragment Y0313-2 were grown at room temperature by vapor diffusion using the hanging drop method. Crystallization buffer containing 0.1 M sodium chloride, 20 mM Tris at pH 7.5, and the VEGF:Fab complex at a concentration of 8 mg/ml was mixed with an equal amount of reservoir solution (15% PEG 4000, 5% isopropanol, 0.1M MES, pH 6.0, 0.2 M Citrate, 0.2 M Ammonium sulfate and 1 mM SPADNS (2-(p-sulfophenylazo)-1,8-dihydroxy-3,6-naphthalene disulphonic acid)). The resulting crystals belong to the monoclinic space group P2 with cell parameters of $a=107.6$ Å, $b=65.8$ Å, $c=123.8$ Å, and $\beta=93.4^\circ$ and contain one VEGF-dimer bound to two Fab fragments in the asymmetric unit.

Prior to flash cooling with liquid nitrogen, crystals were dipped into artificial mother liquor containing 20% glycerol. One diffraction data set was collected from a single crystal at 100 K on a CCD detector at the Advanced Light Source (Berkeley, CA). The data were processed using MOSFLM (Leslie, A *MOSFLM Users Guide*, MRC-LMB, Cambridge (1994)) and programs of the CCP4 suite (Collaborative Computing Project No. 4 *Acta Crystalllog. sect. D*, 50: 760-763 (1994)). The final data set was of good quality ($R_{\text{sym}} = 7.4$ %) with a completeness of 94.5 % for all reflections between 25 Å and 2.8 Å resolution.

Initial phases for the complex were obtained by molecular replacement, using the constant domains and the variable domains of the Fab fragment F(ab)-12 as separate search models. A model of the receptor binding domain of VEGF could be placed unambiguously in a resulting difference density map.

Refinement of the model with program X-PLOR (Bruenger et al. *Science* 235: 458-460 (1987)) resulted in a final R-value of 21.2 % with an R-free of 26.6% using all data between 2.8 Å and 25 Å.

New Antibody-Antigen Contacts in the Insert-Fab Complex with VEGF: The results of x-ray crystallography show that the introduction of the insert (Asn 104a, Glu 104b and Arg 104c (note: numbering of Y0313-2 residues is sequential with inserted residues given a letter, rather than according to Kabat et al., supra) together with the two substitutions

5 (G104V and S105K) enclosing it, increases the total amount of buried
surface in the interface between VEGF and the antibody by about 20% (see
Figure 4), as compared with the structure of the F(ab)-12 complex (Muller
10 *et al.*, *Structure* 6(9):1153-1167 (1998)). The main contributors for the
5 enlargement of the contact area are residues Val 104 and Arg 104c.
Together, these two residues account for additional 220 Å² of buried
surface on the Fab fragment. The side chain of Val 104 is packing tightly
15 against the main chain of residues 93 to 95 of VEGF. The newly introduced
Arg 104c forms a charged interaction with the carboxyl group of Asp 41 of
10 VEGF and is also in contact with the phenyl ring of Tyr 39 (see Figure 5).
Minor contributions to the interface are made by the side chain of Lys 105
which is in the vicinity of the VEGF residues Glu 44 and Tyr 45. The side
20 chains of residues Asn 104a and Glu 104b are pointing away from the
interface and neither of them contributes significantly to the interface
15 between the Fab fragment and VEGF.

5

WHAT IS CLAIMED IS:

10

1. An antibody variant of a parent antibody, which antibody variant comprises an amino acid insertion in or adjacent to a hypervariable region of the parent antibody and has a binding affinity for a target antigen which is at least about two fold stronger than the binding affinity of the parent antibody for said antigen.

15

2. The antibody variant of claim 1 which has an amino acid insertion in a hypervariable region of the parent antibody.

10

20

3. The antibody variant of claim 1 wherein the hypervariable region is Complementarity Determining Region (CDR) H3 of a heavy chain variable domain of the parent antibody.

25

4. The antibody variant of claim 1 wherein about one to about 30 amino acid residues have been inserted in or adjacent to the hypervariable region of the parent antibody.

30

5. The antibody variant of claim 4 wherein about two to about ten amino acid residues have been inserted in or adjacent to the hypervariable region of the parent antibody.

35

6. The antibody variant of claim 1 which has a binding affinity for said antigen that is at least about five fold stronger than the binding affinity of the parent antibody for said antigen.

40

7. The antibody variant of claim 1 wherein the antibody variant has a potency in a biological activity assay which is at least about 20 fold greater than the potency of the parent antibody in the biological activity assay.

45

8. The antibody variant of claim 7 wherein the potency of the antibody variant in the biological activity assay is at least about 50 fold greater than the potency of the parent antibody in the biological activity assay.

35

50

9. The antibody variant of claim 1 wherein the parent antibody is a humanized antibody.

55

5

10. The antibody variant of claim 1 wherein the parent antibody is a human antibody.

10

11. The antibody variant of claim 1 wherein at least one of the inserted residues has a net positive charge or a net negative charge.

15

12. The antibody variant of claim 11 wherein at least one of the inserted residues is arginine or lysine.

10

13. The antibody variant of claim 3 wherein the insertion is adjacent to residue number 100 of the heavy chain variable domain of the parent antibody, utilizing the variable domain residue numbering as in Kabat.

20

14. The antibody variant of claim 13 wherein the insertion consists of about three inserted amino acid residues.

25

15. The antibody variant of claim 1 further comprising an amino acid substitution in the hypervariable region.

30

16. The antibody variant of claim 1 which comprises a heavy chain variable domain, wherein CDR H3 of a heavy chain variable domain of the variant antibody comprises the amino acid sequence of SEQ ID NO:85.

35

17. The antibody variant of claim 15 which comprises a heavy chain variable domain comprising the amino acid sequence in SEQ ID NO:98 or SEQ ID NO:99.

40

18. A composition comprising the antibody variant of claim 1 and a pharmaceutically acceptable carrier.

30

19. An antibody variant comprising a heavy chain variable domain, wherein CDR H3 of the heavy chain variable domain comprises the amino acid sequence of CDR H3 of a variant selected from the group consisting of Y0239-19 (SEQ ID NO:85); Y0239-8 (SEQ ID NO:53); Y0240-1 (SEQ ID NO:86); Y0239-12 (SEQ ID NO:78); Y0239-9 (SEQ ID NO:54); and Y0261-6 (SEQ ID NO:89).

45

35

50

55

5

10

20. A method for producing an antibody variant comprising introducing an amino acid residue in or adjacent to a hypervariable region of a parent antibody, wherein the antibody variant has a binding affinity for a target antigen which is at least about two fold stronger than the binding affinity of the parent antibody for said antigen.

15

21. The method of claim 20 wherein the hypervariable region in which the amino acid residue is introduced is one which is involved in binding the antigen in the parent antibody.

10

20

22. A method for making an antibody variant, comprising the steps of:

(a) identifying potential amino acid interactions between a hypervariable region of a parent antibody and a target antigen;

(b) preparing a variant of the parent antibody comprising introducing an amino acid residue in or adjacent to the hypervariable region of the parent antibody, wherein the introduced amino acid residue contributes to the potential amino acid interactions in (a); and

25

(c) selecting an antibody variant prepared as in (b) which has a stronger binding affinity for said antigen than the parent antibody.

30

20

23. The method of claim 22, wherein step (a) involves analyzing a molecular model of the parent antibody complexed with said antigen.

35

25

24. The method of claim 22 wherein step (b) comprises preparing antibody variants displayed on phage.

40

30

25. The method of claim 22 wherein the amino acid interactions are selected from the group consisting of hydrogen-bonding, Van der Waals interactions and ionic interactions.

45

27. A vector comprising the nucleic acid of claim 26.

35

28. A host cell transformed with the vector of claim 27.

50

29. A process of producing an antibody variant comprising culturing the

55

5

host cell of claim 28 so that the nucleic acid is expressed.

10

30. The process of claim 29 further comprising recovering the antibody variant from the host cell culture.

5

31. The process of claim 30 wherein the antibody variant is recovered from the host cell culture medium.

15

20

25

30

35

40

45

50

55

1 / 6

□ = differences from F(ab)-12

¹⁰ ²⁰ ³⁰
 F(ab)-12 DIQMTQSPSSLSASVGDRVTITCSASQDISNYLNWYQQ
 Y0192 DIQ□TQSPSSLSASVGDRVTITC□RANEQLSNYLNWYQQ
 Y0238-3 DIQ□TQSPSSLSASVGDRVTITC□RANEQLSNYLNWYQQ
 Y0239-19 DIQ□TQSPSSLSASVGDRVTITC□RANEQLSNYLNWYQQ
 Y0313-2 DIQ□TQSPSSLSASVGDRVTITC□RANEQLSNYLNWYQQ

⁴⁰ ⁵⁰ ⁶⁰ ⁷⁰
 F(ab)-12 KPGKAPKVLIIYFTSSLHSGVPSRFSGSGSGTDFTLTIS
 Y0192 KPGKAPKVLIIYFTSSLHSGVPSRFSGSGSGTDFTLTIS
 Y0238-3 KPGKAPKVLIIYFTSSLHSGVPSRFSGSGSGTDFTLTIS
 Y0239-19 KPGKAPKVLIIYFTSSLHSGVPSRFSGSGSGTDFTLTIS
 Y0313-2 KPGKAPKVLIIYFTSSLHSGVPSRFSGSGSGTDFTLTIS

CDR-L1

⁸⁰ ⁹⁰ ¹⁰⁰
 F(ab)-12 SLQPEDFATYYCQQYSTVPWTFGQGTKVEIKRTV (SEQ ID NO:94)
 Y0192 SLQPEDFATYYCQQYSTVPWTFGQGTKVEIKRTV (SEQ ID NO:95)
 Y0238-3 SLQPEDFATYYCQQYSTVPWTFGQGTKVEIKRTV (SEQ ID NO:95)
 Y0239-19 SLQPEDFATYYCQQYSTVPWTFGQGTKVEIKRTV (SEQ ID NO:95)
 Y0313-2 SLQPEDFATYYCQQYSTVPWTFGQGTKVEIKRTV (SEQ ID NO:95)

CDR-L3

FIG. 1A

2 / 6

□ = differences from F(ab)-12

¹⁰ ²⁰ ³⁰
 F(ab)-12 EVQLVESGGGLVQPGGSLRLSCAASGYTF³⁰TNYGMNWVR
 Y0192 EVQLVESGGGLVQPGGSLRLSCAASGYTF³⁰TNYGMNWVR
 Y0238-3 EVQLVESGGGLVQPGGSLRLSCAASGY³⁰TF³⁰HYGMNWVR
 Y0239-19 EVQLVESGGGLVQPGGSLRLSCAASGYTF³⁰TNYGMNWVR
 Y0313-2 EVQLVESGGGLVQPGGSLRLSCAASGY³⁰TF³⁰HYGMNWVR

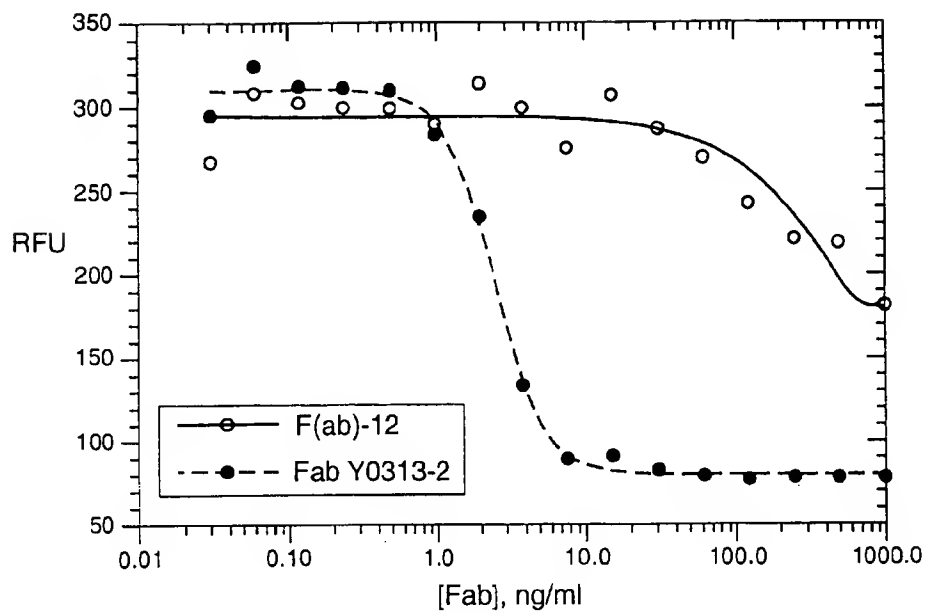
⁴⁰ ⁵⁰ ⁶⁰ ⁷⁰
 F(ab)-12 QAPGKGLEWVGWINTYTGEPTYAADF⁶⁰KRRFT⁷⁰FSLDTSKSTA
 Y0192 QAPGKGLEWVGWINTYTGEPTYAADF⁶⁰KRRFT⁷⁰FSLDTSKSTA
 Y0238-3 QAPGKGLEWVGWINTYTGEPTYAADF⁶⁰KRRFT⁷⁰FSLDTSKSTA
 Y0239-19 QAPGKGLEWVGWINTYTGEPTYAADF⁶⁰KRRFT⁷⁰FSLDTSKSTA
 Y0313-2 QAPGKGLEWVGWINTYTGEPTYAADF⁶⁰KRRFT⁷⁰FSLDTSKSTA

⁸⁰ ⁹⁰ ¹⁰⁰
 F(ab)-12 YLQMNSLRAEDTAVYCAKYPHYG⁹⁰---SSHWYFDVWGQGT¹⁰⁰L (SEQ ID NO:96)
 Y0192 YLQMNSLRAEDTAVYCAKYPHYG⁹⁰---SSHWYFDVWGQGT¹⁰⁰L (SEQ ID NO:96)
 Y0238-3 YLQMNSLRAEDTAVYCAKYPHYG⁹⁰---SSHWYFDVWGQGT¹⁰⁰L (SEQ ID NO:97)
 Y0239-19 YLQMNSLRAEDTAVYCAKYPHY⁹⁰VNERK⁹⁵SHWYFDVWGQGT¹⁰⁰L (SEQ ID NO:98)
 Y0313-2 YLQMNSLRAEDTAVYCAKYPHY⁹⁰VNERK⁹⁵SHWYFDVWGQGT¹⁰⁰L (SEQ ID NO:99)

 CDR-H1 CDR-H2 CDR-H3
 CDR-7

FIG. 1B

3 / 6

**FIG._2**

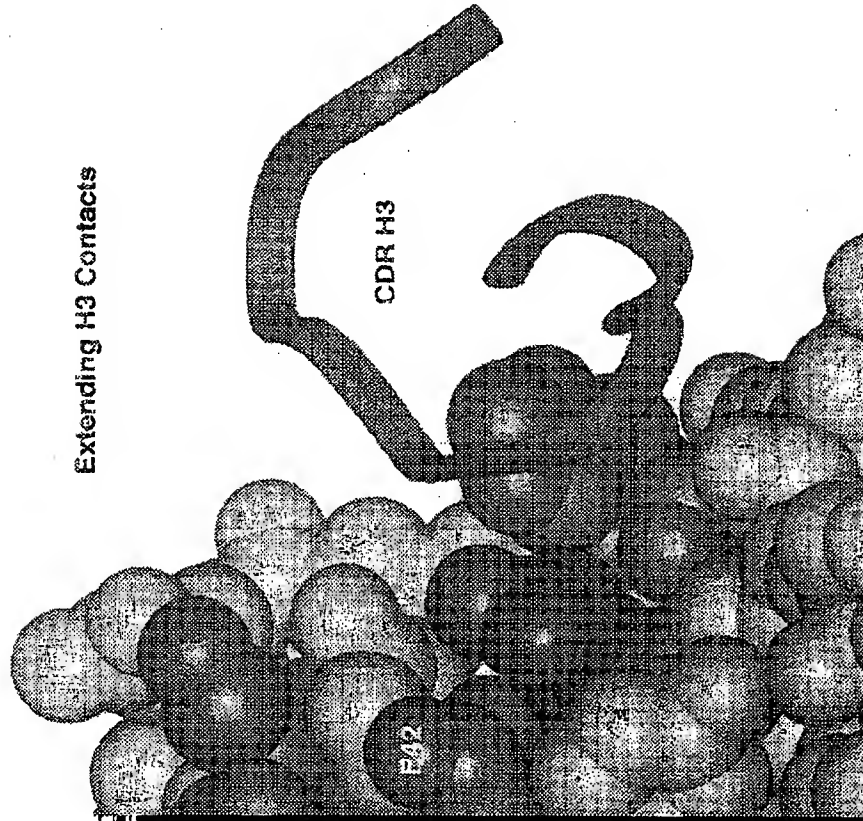


FIG._3

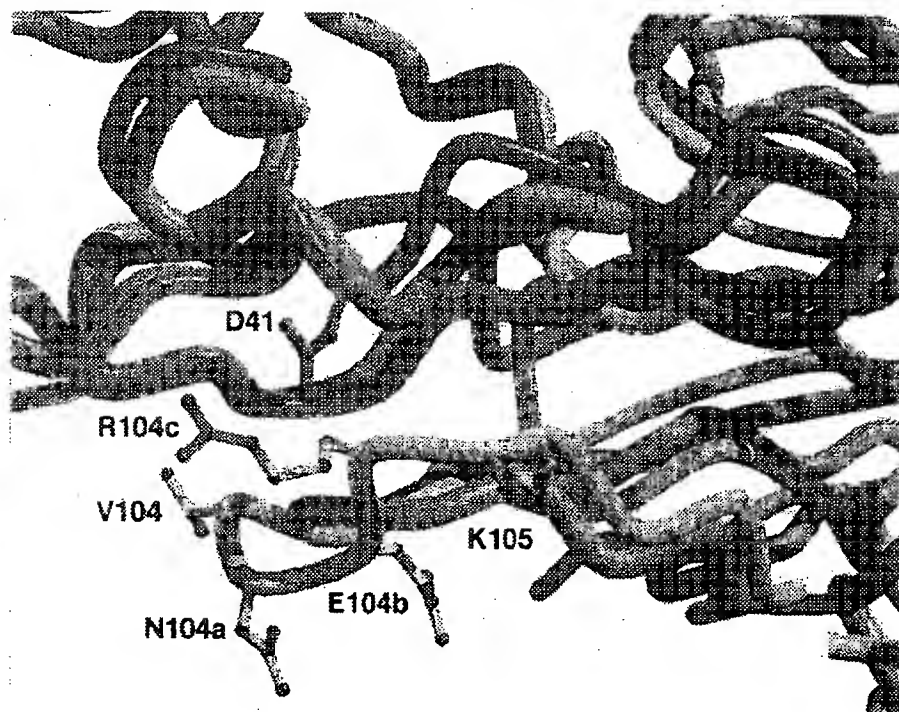


FIG. 4

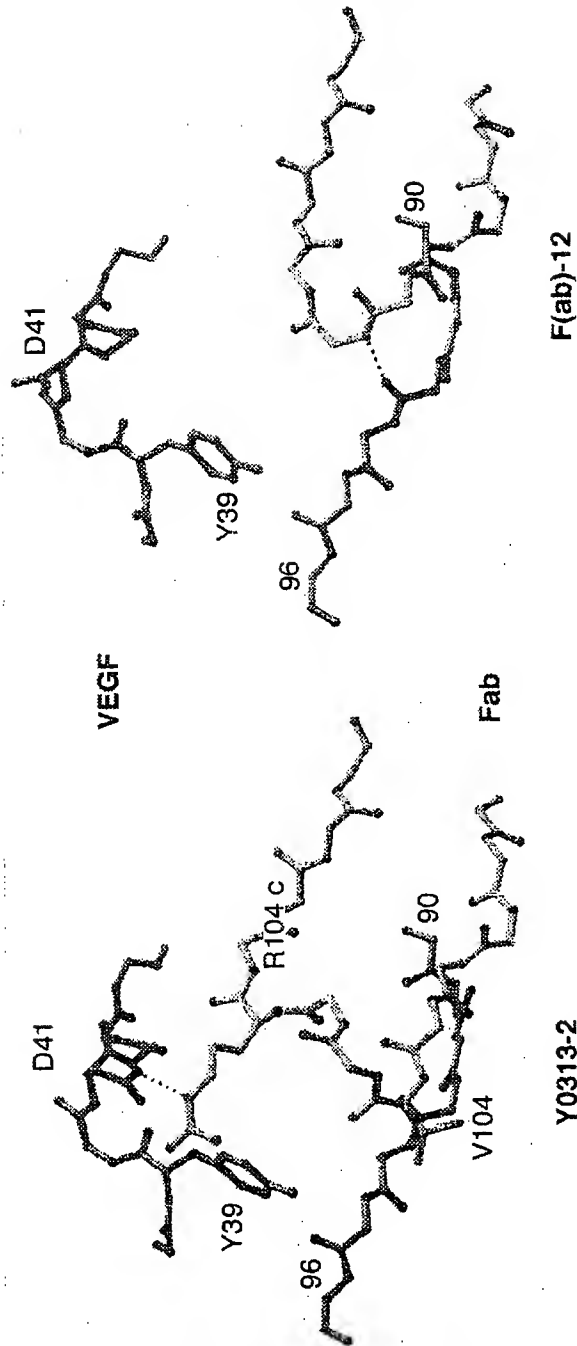


FIG._5

Sequence Listing

<110> Genentech, Inc.

5 <120> ANTIBODY VARIANTS

<130> P1469R1PCT

<141> 1999-11-16

10 <150> US 60/108,945

<151> 1998-11-18

<160> 99

15 <210> 1

<211> 32

<212> DNA

<213> artificial sequence

20 <220>

<221> artificial

<222> 1-32

<223> frameshift oligo

25 <400> 1

ctgtcaacag tataqctacc gtgccg-gga cg 32

<210> 2

30 <211> 32

<212> DNA

<213> artificial sequence

<220>

35 <221> artificial

<222> 1-32

<223> frameshift oligo

<400> 2

40 gcagcttctg gctatgacct tcaccaacta tg 32

<210> 3

<211> 33

<212> DNA

45 <213> artificial sequence

<220>

<221> artificial

<222> 1-33

50 <223> frameshift oligo

<400> 3

gatggattaa cacctatgac cggatgaaccg acc 33

55 <210> 4

<211> 34

<212> DNA

<213> artificial sequence

<220>
<221> artificial
<222> 1-34
<223> frameshift oligo
5
<400> 4
gatggattaa cacctattga accgacctat gctg 34
10
<210> 5
<211> 35
<212> DNA
<213> artificial sequence
15
<220>
<221> artificial
<222> 1-35
<223> frameshift oligo
20
<400> 5
gtacccgcac tattatgagc agccactggg atttc 35
25
<210> 6
<211> 32
<212> DNA
<213> artificial sequence
30
<220>
<221> artificial
<222> 1-32
<223> frameshift oligo
35
<400> 6
gtacccgcac tattatgagc cactgggtatt tc 32
40
<210> 7
<211> 43
<212> DNA
<213> artificial sequence
45
<220>
<221> artificial
<222> 1-43
<223> random oligo
50
<220>
<221> unknown
<222> 17-18, 20-21, 23-24, 26-27
<223> unknown base
55
<400> 7
ctgtcaacag tatagcnnsn nsnnnsnsac cgtgccgtgg acg 43
60
<210> 8
<211> 49
<212> DNA
<213> artificial sequence
65
<220>
<221> artificial

<222> 1-49
<223> random oligo

<220>
5 <221> unknown
<222> 17-18, 20-21, 23-24, 26-27, 29-30, 32-33
<223> unknown base

<400> 8
10 ctgtcaacag tatagcnnsn nsnnnsnnsnn snnsaccgtg ccgtggacg 49

<210> 9
<211> 43
<212> DNA
15 <213> artificial sequence

<220>
<221> artificial
<222> 1-43
20 <223> random oligo

<220>
<221> unknown
<222> 16-17, 19-20, 22-23, 25-26
25 <223> unknown base

<400> 9
gcagcttctg gctatnnsnn snnsnnsacc ttcaccaact atg 43

<210> 10
30 <211> 46
<212> DNA
<213> artificial sequence

<220>
35 <221> artificial
<222> 1-46
<223> random oligo

<220>
40 <221> unknown
<222> 16-17, 19-20, 22-23, 25-26, 28-29
<223> unknown base

<400> 10
45 gcagcttctg gctatnnsnn snnsnnsnns accttcacca actatg 46

<210> 11
<211> 49
50 <212> DNA
<213> artificial sequence

<220>
55 <221> artificial
<222> 1-49
<223> random oligo

<220>
<221> unknown

<222> 16-17, 19-20, 22-23, 25-26, 28-29, 31-32
 <223> unknown base

5 <400> 11
 gcagcttctg gctatnnsnn snnsnnsnns nnsaccttca ccaactatg 49

<210> 12
 <211> 41
 <212> DNA
 10 <213> artificial sequence

<220>
 <221> artificial
 <222> 1-41
 15 <223> random oligo

<220>
 <221> unknown
 <222> 18-19, 21-22, 24-25
 20 <223> unknown base

<400> 12
 gatggattaa cacctatnns nnsnnsaccg gtgaaccgac c 41

25 <210> 13
 <211> 44
 <212> DNA
 <213> artificial sequence

30 <220>
 <221> artificial
 <222> 1-44
 <223> random oligo

35 <220>
 <221> unknown
 <222> 18-19, 21-22, 24-25, 27-28
 <223> unknown base

40 <400> 13
 gatggattaa cacctatnns nnsnnsnnsa ccggtgaacc gacc 44

<210> 14
 <211> 51
 45 <212> DNA
 <213> artificial sequence

<220>
 <221> artificial
 50 <222> 1-51
 <223> random oligo

<220>
 <221> unknown
 55 <222> 18-19, 21-22, 24-25, 27-28, 30-31, 33-34
 <223> unknown base

<400> 14
 gatggattaa cacctatnns nnsnnsnnsn nsnnsgaacc gacctatgct 50

g 51

5 <210> 15
<211> 46
<212> DNA
<213> artificial sequence

10 <220>
<221> artificial
<222> 1-46
<223> random oligo

15 <220>
<221> unknown
<222> 17-18, 20-21, 23-24, 26-27
<223> unknown base

20 <400> 15
gtaccgcac tattatnnsn nsnnnsnsag cagccactgg tatttc 46

25 <210> 16
<211> 46
<212> DNA
<213> artificial sequence

30 <220>
<221> artificial
<222> 1-46
<223> random oligo

35 <220>
<221> unknown
<222> 17-18, 20-21, 23-24, 26-27
<223> unknown base

40 <400> 16
gtaccgcac tattatnnsn nsnnnsnsag cagccactgg tatttc 46

45 <210> 17
<211> 46
<212> DNA
<213> artificial sequence

50 <220>
<221> artificial
<222> 1-46
<223> random oligo

55 <220>
<221> unknown
<222> 17-18, 20-21, 23-24, 26-27, 29-30
<223> unknown base

<400> 17
gtaccgcac tattatnnsn nsnnnsnsnn sagccactgg tatttc 46

<210> 18
<211> 49

```

<212> DNA
<213> artificial sequence

<220>
5  <221> artificial
   <222> 1-49
   <223> random oligo

<220>
10 <221> unknown
    <222> 17-18, 20-21, 23-24, 26-27, 29-30, 32-33
    <223> unknown base

<400> 18
15  gtaccgcgcac tattatnnsh nsnrshnnsh snnsagccac tggatttc 49

<210> 19
<211> 10
<212> PRT
20 <213> artificial sequence

<220>
<221> artificial
<222> 1-10
25 <223> variant CDR sequence

<400> 19
    Gly Tyr Asp Phe Thr Asn Tyr Gly Ile Asn
      1             5             10

30 <210> 20
    <211> 10
    <212> PRT
    <213> artificial sequence
35 <220>
    <221> artificial
    <222> 1-10
    <223> variant CDR sequence
40 <400> 20
    Gly Tyr Asp Tyr Thr Asn Tyr Gly Ile Asn
      1             5             10

45 <210> 21
    <211> 10
    <212> PRT
    <213> artificial sequence

50 <220>
    <221> artificial
    <222> 1-10
    <223> variant CDR sequence

55 <400> 21
    Gly Tyr Asp Trp Thr Asn Tyr Gly Ile Asn
      1             5             10

<210> 22

```

<211> 13
<212> PRT
<213> artificial sequence

5 <220>
<221> artificial
<222> 1-13
<223> variant CDR sequence

10 <400> 22
Trp Ile Asn Thr Trp Thr Gly Glu Pro Thr Tyr Ala Ala
1 5 10 13

<210> 23
15 <211> 9
<212> PRT
<213> artificial sequence

<220>
20 <221> artificial
<222> 1-9
<223> variant CDR sequence

<400> 23
25 Gln Gln Tyr Ser Ala Thr Pro Trp Thr
1 5 9

<210> 24
30 <211> 9
<212> PRT
<213> artificial sequence

<220>
35 <221> artificial
<222> 1-9
<223> variant CDR sequence

<400> 24
40 Gln Gln Tyr Ser Asn Val Pro Trp Thr
1 5 9

<210> 25
45 <211> 9
<212> PRT
<213> artificial sequence

<220>
50 <221> artificial
<222> 1-9
<223> variant CDR sequence

<400> 25
55 Gln Gln Tyr Ser Ala Val Pro Trp Thr
1 5 9

<210> 26
<211> 9
<212> PRT
<213> artificial sequence

```

<220>
<221> artificial
<222> 1-9
5 <223> variant CDR sequence

<400> 26
  Gln Gln Tyr Ser Ser Val Pro Trp Thr
    1             5             9
10
<210> 27
<211> 17
<212> PRT
<213> artificial sequence
15
<220>
<221> artificial
<222> 1-17
<223> variant CDR sequence
20
<400> 27
  Tyr Pro His Tyr Tyr Ala Lys Glu Arg Ser Ser His Trp Tyr Phe
    1             5             10             15
25  Asp Val
    17

<210> 28
<211> 17
30 <212> PRT
<213> artificial sequence

<220>
<221> artificial
35 <222> 1-17
<223> variant CDR sequence

<400> 28
  Tyr Pro His Tyr Tyr Val Gly Glu Thr Ser Ser His Trp Tyr Phe
    1             5             10             15
40  Asp Val
    17

45 <210> 29
<211> 17
<212> PRT
<213> artificial sequence

50 <220>
<221> artificial
<222> 1-17
<223> variant CDR sequence

55 <400> 29
  Tyr Pro His Tyr Tyr Ala Arg Asp Arg Ser Ser His Trp Tyr Phe
    1             5             10             15

  Asp Val

```

```

17
<210> 30
<211> 18
5  <212> PRT
   <213> artificial sequence

<220>
<221> artificial
10 <222> 1-18
   <223> variant CDR sequence

<400> 30
15 Tyr Pro His Tyr Tyr Glu Arg Asp Gly Lys Ser Ser His Trp Tyr
    1             5             10             15

    Phe Asp Val
      18

20 <210> 31
   <211> 17
   <212> PRT
   <213> artificial sequence

25 <220>
   <221> artificial
   <222> 1-17
   <223> variant CDR sequence

30 <400> 31
   Tyr Pro His Tyr Tyr Arg Asn Glu Lys Ser Ser His Trp Tyr Phe
    1             5             10             15

    Asp Val
      17

35 <210> 32
   <211> 17
   <212> PRT
40 <213> artificial sequence

   <220>
   <221> artificial
   <222> 1-17
45 <223> variant CDR sequence

   <400> 32
   Tyr Pro His Tyr Tyr Val Gly Glu Gln Ser Ser His Trp Tyr Phe
    1             5             10             15

50   Asp Val
      17

55 <210> 33
   <211> 17
   <212> PRT
   <213> artificial sequece

   <220>

```



```

<221> artificial
<222> 1-17
<223> variant CDR sequence

5  <400> 33
    Tyr Pro His Tyr Tyr Gln Arg Asp Arg Ser Ser His Trp Tyr Phe
      1             5             10             15

    Asp Val
10      17

    <210> 34
    <211> 18
    <212> PRT
15  <213> artificial sequence

    <220>
    <221> artificial
    <222> 1-18
20  <223> variant CDR sequence

    <400> 34
    Tyr Pro His Tyr Tyr Gln Lys Gln Ser Lys Ser Ser His Trp Tyr
      1             5             10             15

25  Phe Asp Val
      18

    <210> 35
    <211> 18
    <212> PRT
30  <213> artificial sequence

    <220>
    <221> artificial
    <222> 1-18
35  <223> variant CDR sequence

    <400> 35
40  Tyr Pro His Tyr Tyr Gln Asn Glu Gly Pro Ser Ser His Trp Tyr
      1             5             10             15

    Phe Asp Val
      18

45  <210> 36
    <211> 17
    <212> PRT
    <213> artificial sequence

50  <220>
    <221> artificial
    <222> 1-17
    <223> variant CDR sequence

55  <400> 36
    Tyr Pro His Tyr Tyr Gly Asn His Arg Ser Ser His Trp Tyr Phe
      1             5             10             15

```

```

      Asp Val
      17

5    <210> 37
    <211> 17
    <212> PRT
    <213> artificial sequence

    <220>
10   <221> artificial
    <222> 1-17
    <223> variant CDR sequence

    <400> 37
15   Tyr Pro His Tyr Tyr Arg Thr Glu Lys Ser Ser His Trp Tyr Phe
      1             5             10             15

      Asp Val
      17

20   <210> 38
    <211> 17
    <212> PRT
    <213> artificial sequence

25   <220>
    <221> artificial
    <222> 1-17
    <223> variant CDR sequence

30   <400> 38
    Tyr Pro His Tyr Tyr Leu Lys Asp Arg Ser Ser His Trp Tyr Phe
      1             5             10             15

35   Asp Val
      17

    <210> 39
    <211> 17
40   <212> PRT
    <213> artificial sequence

    <220>
    <221> artificial
45   <222> 1-17
    <223> variant CDR sequence

    <400> 39
50   Tyr Pro His Tyr Tyr Gln Asp Glu Lys Ser Ser His Trp Tyr Phe
      1             5             10             15

      Asp Val
      17

55   <210> 40
    <211> 17
    <212> PRT
    <213> artificial sequence

```

```

<220>
<221> artificial
<222> 1-17
<223> variant CDR sequence
5
<400> 40
Tyr Pro His Tyr Tyr Val Gly Glu Lys Ser Ser His Trp Tyr Phe
1 5 10 15

10 Asp Val
17

<210> 41
<211> 17
15 <212> PRT
<213> artificial sequence

<220>
<221> artificial
20 <222> 1-17
<223> variant CDR sequence

<400> 41
Tyr Pro His Tyr Tyr Arg Asp Glu Arg Ser Ser His Trp Tyr Phe
25 1 5 10 15

Asp Val
17

30 <210> 42
<211> 17
<212> PRT
<213> artificial sequence

35 <220>
<221> artificial
<222> 1-17
<223> variant CDR sequence

40 <400> 42
Tyr Pro His Tyr Tyr Thr Tyr Asp Lys Ser Ser His Trp Tyr Phe
1 5 10 15

Asp Val
45 17

<210> 43
<211> 18
<212> PRT
50 <213> artificial sequence

<220>
<221> artificial
<222> 1-18
55 <223> variant CDR sequence

<400> 43
Tyr Pro His Tyr Tyr His Thr Arg Gly Gly Ser Ser His Trp Tyr
1 5 10 15

```

Phe Asp Val
18

5 <210> 44
<211> 17
<212> PRT
<213> artificial sequence

10 <220>
<221> artificial
<222> 1-17
<223> variant CDR sequence

15 <400> 44
Tyr Pro His Tyr Tyr Leu Asn Asp Lys Ser Ser His Trp Tyr Phe
1 5 10 15

Asp Val
17

20 <210> 45
<211> 17
<212> PRT
<213> artificial sequence

25 <220>
<221> artificial
<222> 1-17
<223> variant CDR sequence

30 <400> 45
Tyr Pro His Tyr Tyr Arg Asp Arg Ser Ser His Trp Tyr Phe
1 5 10 15

35 Asp Val
17

40 <210> 46
<211> 17
<212> PRT
<213> artificial sequence

45 <220>
<221> artificial
<222> 1-17
<223> variant CDR sequence

50 <400> 46
Tyr Pro His Tyr Tyr Arg Asn Glu Arg Ser Ser His Trp Tyr Phe
1 5 10 15

Asp Val
17

55 <210> 47
<211> 17
<212> PRT
<213> artificial sequence

```

<220>
<221> artificial
<222> 1-17
5 <223> variant CDR sequence

<400> 47
  Tyr Pro His Tyr Tyr Lys Asn Asp Lys Ser Ser His Trp Tyr Phe
    1             5             10             15
10   Asp Val
    17

<210> 48
15 <211> 17
    <212> PRT
    <213> artificial sequence

<220>
20 <221> artificial
    <222> 1-17
    <223> variant CDR sequence

<400> 48
25   Tyr Pro His Tyr Tyr Leu Ala Asp Arg Ser Ser His Trp Tyr Phe
    1             5             10             15

    Asp Val
    17
30

<210> 49
<211> 17
<212> PRT
<213> artificial sequence
35

<220>
<221> artificial
<222> 1-17
<223> variant CDR sequence
40

<400> 49
  Tyr Pro His Tyr Tyr Val Asn Glu Arg Ser Ser His Trp Tyr Phe
    1             5             10             15

45   Asp Val
    17

<210> 50
<211> 17
50 <212> PRT
    <213> artifical sequence

<220>
<221> artificial
55 <222> 1-17
    <223> variant CDR sequence

<400> 50
  Tyr Pro His Tyr Tyr Leu Lys Asp Lys Ser Ser His Trp Tyr Phe

```

```

      1             5             10             15

      Asp Val
      17
5
      <210> 51
      <211> 18
      <212> PRT
      <213> artificial sequence
10
      <220>
      <221> artificial
      <222> 1-18
      <223> variant CDR sequence
15
      <400> 51
      Tyr Pro His Tyr Tyr Leu Lys Asp Gly Arg Ser Ser His Trp Tyr
      1             5             10             15

20      Phe Asp Val
      18

      <210> 52
      <211> 18
25      <212> PRT
      <213> artificial sequence

      <220>
      <221> artificial
30      <222> 1-18
      <223> variant CDR sequence

      <400> 52
      Tyr Pro His Tyr Tyr Glu Arg Asp Gly Arg Ser Ser His Trp Tyr
35      1             5             10             15

      Phe Asp Val
      18

40      <210> 53
      <211> 18
      <212> PRT
      <213> artificial sequence

45      <220>
      <221> artificial
      <222> 1-18
      <223> variant CDR sequence

50      <400> 53
      Tyr Pro His Tyr Tyr Leu Arg Asp Gly Arg Ser Ser His Trp Tyr
      1             5             10             15

      Phe Asp Val
55      18

      <210> 54
      <211> 16
      <212> PRT

```

<213> artificial sequence
 <220>
 <221> artificial
 5 <222> 1-16
 <223> variant CDR sequence
 <400> 54
 Tyr Pro His Tyr Tyr Leu Gly Glu Ser Ser His Trp Tyr Phe Asp
 10 1 5 10 15
 Val
 16
 15 <210> 55
 <211> 17
 <212> PRT
 <213> artificial sequence
 20 <220>
 <221> artificial
 <222> 1-17
 <223> variant CDR sequence
 25 <400> 55
 Tyr Pro His Tyr Tyr Leu Gly Glu Lys Ser Ser His Trp Tyr Phe
 1 5 10 15
 Asp Val
 30 17
 <210> 56
 <211> 18
 <212> PRT
 35 <213> artificial sequence
 <220>
 <221> artificial
 <222> 1-18
 40 <223> variant CDR sequence
 <400> 56
 Tyr Pro His Tyr Tyr Leu Lys Asp Arg Arg Ser Ser His Trp Tyr
 1 5 10 15
 45 Phe Asp Val
 18
 <210> 57
 50 <211> 18
 <212> PRT
 <213> artificial sequence
 <220>
 55 <221> artificial
 <222> 1-18
 <223> variant CDR sequence
 <400> 57

```

Tyr Pro His Tyr Tyr Leu Lys Asp Gly Met Ser Ser His Trp Tyr
  1             5             10             15

5   Phe Asp Val
    18

<210> 58
<211> 17
<212> PRT
10 <213> artificial sequence

<220>
<221> artificial
<222> 1-17
15 <223> variant CDR sequence

<400> 58
Tyr Pro His Tyr Tyr Glu Lys Gln Arg Lys Ser His Trp Tyr Phe
  1             5             10             15

20  Asp Val
    17

<210> 59
25 <211> 17
<212> PRT
<213> artificial sequence

<220>
30 <221> artificial
<222> 1 17
<223> variant CDR sequence

<400> 59
35 Tyr Pro His Tyr Tyr Lys Glu Asp Lys Lys Ser His Trp Tyr Phe
  1             5             10             15

    Asp Val
      17

40 <210> 60
<211> 17
<212> PRT
<213> artificial sequence

45 <220>
<221> artificial
<222> 1-17
<223> variant CDR sequence

50 <400> 60
Tyr Pro His Tyr Tyr Ser His Gln Lys Arg Ser His Trp Tyr Phe
  1             5             10             15

55  Asp Val
    17

<210> 61
<211> 17

```



```

<212> PRT
<213> artificial sequence

<220>
5  <221> artificial
   <222> 1-17
   <223> variant CDR sequence

<400> 61
10  Tyr Pro His Tyr Tyr Ser Gly Glu Arg Glu Ser His Trp Tyr Phe
    1             5             10             15

    Asp Val
    17
15
    <210> 62
    <211> 17
    <212> PRT
    <213> artificial sequence
20
    <220>
    <221> artificial
    <222> 1-17
    <223> variant CDR sequence
25
    <400> 62
    Tyr Pro His Tyr Tyr Gln Ser Glu Gly Arg Ser His Trp Tyr Phe
    1             5             10             15

    Asp Val
    17
30
    <210> 63
    <211> 17
    <212> PRT
    <213> artificial sequence
35
    <220>
    <221> artificial
    <222> 1-17
    <223> variant CDR sequence
40
    <400> 63
    Tyr Pro His Tyr Tyr Ser Val Glu Gly Gly Ser His Trp Tyr Phe
    1             5             10             15

    Asp Val
    17
50
    <210> 64
    <211> 17
    <212> PRT
    <213> artificial sequence
55
    <220>
    <221> artificial
    <222> 1-17
    <223> variant CDR sequence

```

<400> 64
Tyr Pro His Tyr Tyr Pro Ser Pro Arg Gly Ser His Trp Tyr Phe
1 5 10 15

5 Asp Val
17

<210> 65
<211> 17
10 <212> PRT
<213> artificial sequence

<220>
<221> artificial
15 <222> 1-17
<223> variant CDR sequence

<400> 65
Tyr Pro His Tyr Tyr Gln Arg Asn Gly Lys Ser His Trp Tyr Phe
20 1 5 10 15

Asp Val
17

25 <210> 66
<211> 17
<212> PRT
<213> artificial sequence

30 <220>
<221> artificial
<222> 1-17
<223> variant CDR sequence

35 <400> 66
Tyr Pro His Tyr Tyr Ala Arg Glu Gly Gly Ser His Trp Tyr Phe
1 5 10 15

Asp Val
40 17

<210> 67
<211> 17
<212> PRT
45 <213> artificial sequence

<220>
<221> artificial
<222> 1-17
50 <223> variant CDR sequence

<400> 67
Tyr Pro His Tyr Tyr Ser Asn Glu Arg Lys Ser His Trp Tyr Phe
1 5 10 15

55 Asp Val
17

<210> 68

<211> 17
<212> PRT
<213> artificial sequence

5 <220>
<221> artificial
<222> 1-17
<223> variant CDR sequence

10 <400> 68
Tyr Pro His Tyr Tyr Arg Gly Asp Arg Lys Ser His Trp Tyr Phe
1 5 10 15
Asp Val
15 17

<210> 69
<211> 17
<212> PRT
20 <213> artificial sequence

<220>
<221> artificial
<222> 1-17
25 <223> variant CDR sequence

<400> 69
Tyr Pro His Tyr Tyr Ser Asp Glu Lys Lys Ser His Trp Tyr Phe
1 5 10 15
30 Asp Val
17

<210> 70
35 <211> 17
<212> PRT
<213> artificial sequence

<220>
40 <221> artificial
<222> 1-17
<223> variant CDR sequence

<400> 70
45 Tyr Pro His Tyr Tyr Arg Ser Gln Arg Lys Ser His Trp Tyr Phe
1 5 10 15
Asp Val
17

50 <210> 71
<211> 18
<212> PRT
<213> artificial sequence

55 <220>
<221> artificial
<222> 1-18
<223> variant CDR sequence

```

<400> 71
  Tyr Pro His Tyr Tyr Ala Trp Arg Asp Arg Arg Ser His Trp Tyr
    1           5           10           15
5   Phe Asp Val
    18

<210> 72
10 <211> 18
    <212> PRT
    <213> artificial sequence

    <220>
15 <221> artificial
    <222> 1-18
    <223> variant CDR sequence

<400> 72
20  Tyr Pro His Tyr Tyr Ala Asn Arg Glu Arg Lys Ser His Trp Tyr
    1           5           10           15

    Phe Asp Val
    18
25

<210> 73
    <211> 17
    <212> PRT
    <213> artificial sequence
30

    <220>
    <221> artificial
    <222> 1-17
    <223> variant CDR sequence
35

<400> 73
    Tyr Pro His Tyr Tyr Val Asn Asp Lys Thr Ser His Trp Tyr Phe
      1           5           10           15

40  Asp Val
    17

    <210> 74
    <211> 17
45 <212> PRT
    <213> artificial sequence

    <220>
    <221> artificial
50 <222> 1-17
    <223> variant CDR sequence

    <400> 74
55  Tyr Pro His Tyr Tyr Val Glu Glu Thr Glu Ser His Trp Tyr Phe
      1           5           10           15

    Asp Val
    17

```

```

<210> 75
<211> 17
<212> PRT
<213> artificial sequence
5
<220>
<221> artificial
<222> 1-17
<223> variant CDR sequence
10
<400> 75
Tyr Pro His Tyr Tyr Glu Lys Glu Arg Lys Ser His Trp Tyr Phe
    1              5              10              15
15
Asp Val
    17

<210> 76
<211> 17
<212> PRT
<213> artificial sequence
20
<220>
<221> artificial
<222> 1-17
<223> variant CDR sequence
25
<400> 76
Tyr Pro His Tyr Tyr Ser His Glu Arg Val Ser His Trp Tyr Phe
30    1              5              10              15

Asp Val
    17

35
<210> 77
<211> 17
<212> PRT
<213> artificial sequence
40
<220>
<221> artificial
<222> 1-17
<223> variant CDR sequence
45
<400> 77
Tyr Pro His Tyr Tyr Arg Asp Glu Arg Glu Ser His Trp Tyr Phe
    1              5              10              15

Asp Val
50    17

<210> 78
<211> 17
<212> PRT
55
<213> artificial sequence

<220>
<221> artificial
<222> 1-17

```

```

<223> variant CDR sequence

<400> 78
5   Tyr Pro His Tyr Tyr Ala His Glu Lys Lys Ser His Trp Tyr Phe
    1           5           10           15

    Asp Val
      17

10  <210> 79
    <211> 17
    <212> PRT
    <213> artificial sequence

15  <220>
    <221> artificial
    <222> 1-17
    <223> variant CDR sequence

20  <400> 79
    Tyr Pro His Tyr Tyr Leu Lys Asp Arg Lys Ser His Trp Tyr Phe
      1           5           10           15

    Asp Val
25  17

    <210> 80
    <211> 17
    <212> PRT
30  <213> artificial sequence

    <220>
    <221> artificial
    <222> 1-17
35  <223> variant CDR sequence

    <400> 80
    Tyr Pro His Tyr Tyr Gln His Asp Arg Thr Ser His Trp Tyr Phe
      1           5           10           15

40  Asp Val
    17

    <210> 81
45  <211> 17
    <212> PRT
    <213> artificial sequence

    <220>
50  <221> artificial
    <222> 1-17
    <223> variant CDR sequence

    <400> 81
55  Tyr Pro His Tyr Tyr Val Thr Asp Arg Lys Ser His Trp Tyr Phe
      1           5           10           15

    Asp Val
      17

```

<210> 82
<211> 17
<212> PRT
5 <213> artificial sequence

<220>
<221> artificial
<222> 1-17
10 <223> variant CDR sequence

<400> 82
Tyr Pro His Tyr Tyr Leu Arg Asp Lys Lys Ser His Trp Tyr Phe
1 5 10 15
15 Asp Val
17

<210> 83
20 <211> 17
<212> PRT
<213> artificial sequence

<220>
25 <221> artificial
<222> 1-17
<223> variant CDR sequence

<400> 83
30 Tyr Pro His Tyr Tyr Ser His Glu Arg Lys Ser His Trp Tyr Phe
1 5 10 15
Asp Val
17
35

<210> 84
<211> 17
<212> PRT
<213> artificial sequence
40

<220>
<221> artificial
<222> 1-17
45 <223> variant CDR sequence

<400> 84
Tyr Pro His Tyr Tyr Leu Asn Glu Arg Lys Ser His Trp Tyr Phe
1 5 10 15
50 Asp Val
17

<210> 85
55 <211> 17
<212> PRT
<213> artificial sequence

<220>
<221> artificial

<222> 1-17
<223> variant CDR sequence

<400> 85
5 Tyr Pro His Tyr Tyr Val Asn Glu Arg Lys Ser His Trp Tyr Phe
1 5 10 15

Asp Val
17

10 <210> 86
<211> 17
<212> PRT
<213> artificial sequence

15 <220>
<221> artificial
<222> 1-17
<223> variant CDR sequence

20 <400> 86
Tyr Pro His Tyr Tyr Leu Thr Asp His Lys Ser His Trp Tyr Phe
1 5 10 15

25 Asp Val
17

<210> 87
<211> 18
30 <212> PRT
<213> artificial sequence

<220>
<221> artificial
35 <222> 1-18
<223> variant CDR sequence

<400> 87
40 Tyr Pro His Tyr Tyr Leu Lys Asp Gly Lys Lys Ser His Trp Tyr
1 5 10 15

Phe Asp Val
18

45 <210> 88
<211> 17
<212> PRT
<213> artificial sequence

50 <220>
<221> artificial
<222> 1-17
<223> variant CDR sequence

55 <400> 88
Tyr Pro His Tyr Tyr Arg Arg Asp Lys Lys Ser His Trp Tyr Phe
1 5 10 15

Asp Val

17
<210> 89
<211> 17
5 <212> PRT
<213> artificial sequence

<220>
<221> artificial
10 <222> 1-17
<223> variant CDR sequence

<400> 89
Tyr Pro His Tyr Tyr Leu Lys Asp Lys Lys Ser His Trp Tyr Phe
15 1 5 10 15

Asp Val
17
20 <210> 90
<211> 17
<212> PRT
<213> artificial sequence

25 <220>
<221> artificial
<222> 1-17
<223> variant CDR sequence

30 <400> 90
Tyr Pro His Tyr Tyr Leu His Asp Arg Lys Ser His Trp Tyr Phe
1 5 10 15

Asp Val
35 17

<210> 91
<211> 17
<212> PRT
40 <213> artificial sequence

<220>
<221> artificial
<222> 1-17
45 <223> variant CDR sequence

<400> 91
Tyr Pro His Tyr Tyr Leu Ser Asp Lys Lys Ser His Trp Tyr Phe
1 5 10 15
50 Asp Val
17

<210> 92
55 <211> 17
<212> PRT
<213> artificial sequence

<220>

```

<221> artificial
<222> 1-17
<223> variant CDR sequence

5  <400> 92
    Tyr Pro His Tyr Tyr Val Asn Glu Arg Lys Ser His Trp Tyr Phe
      1             5             10             15

    Asp Val
10      17

    <210> 93
    <211> 45
    <212> DNA
15  <213> artificial sequence

    <220>
    <221> artificial
    <222> 1-45
20  <223> mutagenesis oligo

    <400> 93
    taccgcgact attatgtgaa cgagcggaag agccactggt atttc 45

25  <210> 94
    <211> 110
    <212> PRT
    <213> artificial sequence

30  <220>
    <221> artificial
    <222> 1-110
    <223> humanized antibody light chain variable domain

35  <400> 94
    Asp Ile Gln Met Thr Gln Ser Pro Ser Ser Leu Ser Ala Ser Val
      1             5             10             15

    Gly Asp Arg Val Thr Ile Thr Cys Ser Ala Ser Gln Asp Ile Ser
40      20             25             30

    Asn Tyr Leu Asn Trp Tyr Gln Gln Lys Pro Gly Lys Ala Pro Lys
      35             40             45

45  Val Leu Ile Tyr Phe Thr Ser Ser Leu His Ser Gly Val Pro Ser
      50             55             60

    Arg Phe Ser Gly Ser Gly Ser Gly Thr Asp Phe Thr Leu Thr Ile
      65             70             75

50  Ser Ser Leu Gln Pro Glu Asp Phe Ala Thr Tyr Tyr Cys Gln Gln
      80             85             90

    Tyr Ser Thr Val Pro Trp Thr Phe Gly Gln Gly Thr Lys Val Glu
55      95             100            105

    Ile Lys Arg Thr Val
      110

```

```

<210> 95
<211> 110
<212> PRT
<213> artificial sequence
5
<220>
<221> artificial
<222> 1-110
<223> humanized antibody light chain variable domain
10
<400> 95
Asp Ile Gln Leu Thr Gln Ser Pro Ser Ser Leu Ser Ala Ser Val
  1             5             10             15
15 Gly Asp Arg Val Thr Ile Thr Cys Arg Ala Asn Glu Gln Leu Ser
      20             25             30
    Asn Tyr Leu Asn Trp Tyr Gln Gln Lys Pro Gly Lys Ala Pro Lys
      35             40             45
20 Val Leu Ile Tyr Phe Thr Ser Ser Leu His Ser Gly Val Pro Ser
      50             55             60
25 Arg Phe Ser Gly Ser Gly Ser Gly Thr Asp Phe Thr Leu Thr Ile
      65             70             75
    Ser Ser Leu Gln Pro Glu Asp Phe Ala Thr Tyr Tyr Cys Gln Gln
      80             85             90
30 Tyr Ser Thr Val Pro Trp Thr Phe Gly Gln Gly Thr Lys Val Glu
      95            100            105
    Ile Lys Arg Thr Val
      110
35
<210> 96
<211> 118
<212> PRT
<213> artificial sequence
40
<220>
<221> artificial
<222> 1-118
<223> humanized antibody heavy chain variable domain
45
<400> 96
Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly
  1             5             10             15
50 Gly Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Tyr Thr Phe Thr
      20             25             30
    Asn Tyr Gly Met Asn Trp Val Arg Gln Ala Pro Gly Lys Gly Leu
      35             40             45
55 Glu Trp Val Gly Trp Ile Asn Thr Tyr Thr Gly Glu Pro Thr Tyr
      50             55             60
    Ala Ala Asp Phe Lys Arg Arg Phe Thr Phe Ser Leu Asp Thr Ser

```

```

        65              70              75
Lys Ser Thr Ala Tyr Leu Gln Met Asn Ser Leu Arg Ala Glu Asp
        80              85              90
5  Thr Ala Val Tyr Tyr Cys Ala Lys Tyr Pro His Tyr Tyr Gly Ser
        95              100             105
10 Ser His Trp Tyr Phe Asp Val Trp Gly Gln Gly Thr Leu
        110             115             118
<210> 97
<211> 118
<212> PRT
15 <213> artificial sequence

<220>
<221> artificial
<222> 1-118
20 <223> humanized antibody heavy chain variable domain

<400> 97
Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly
  1           5           10           15
25 Gly Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Tyr Asp Phe Thr
        20           25           30
30 His Tyr Gly Met Asn Trp Val Arg Gln Ala Pro Gly Lys Gly Leu
        35           40           45
    Glu Trp Val Gly Trp Ile Asn Thr Tyr Thr Gly Glu Pro Thr Tyr
        50           55           60
35 Ala Ala Asp Phe Lys Arg Arg Phe Thr Phe Ser Leu Asp Thr Ser
        65           70           75
    Lys Ser Thr Ala Tyr Leu Gln Met Asn Ser Leu Arg Ala Glu Asp
        80           85           90
40 Thr Ala Val Tyr Tyr Cys Ala Lys Tyr Pro His Tyr Tyr Gly Ser
        95           100          105
45 Ser His Trp Tyr Phe Asp Val Trp Gly Gln Gly Thr Leu
        110          115          118
<210> 98
<211> 121
<212> PRT
50 <213> artificial sequence

<220>
<221> artificial
<222> 1-121
55 <223> humanized antibody heavy chain variable domain

<400> 98
Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly
  1           5           10           15

```

```

      Gly Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Tyr Thr Phe Thr
      20                               25                               30

5      Asn Tyr Gly Met Asn Trp Val Arg Gln Ala Pro Gly Lys Gly Leu
      35                               40                               45

      Glu Trp Val Gly Trp Ile Asn Thr Tyr Thr Gly Glu Pro Thr Tyr
      50                               55                               60

10     Ala Ala Asp Phe Lys Arg Arg Phe Thr Phe Ser Leu Asp Thr Ser
      65                               70                               75

      Lys Ser Thr Ala Tyr Leu Gln Met Asn Ser Leu Arg Ala Glu Asp
15     80                               85                               90

      Thr Ala Val Tyr Tyr Cys Ala Lys Tyr Pro His Tyr Tyr Val Asn
      95                               100                              105

20     Glu Arg Lys Ser His Trp Tyr Phe Asp Val Trp Gly Gln Gly Thr
      110                              115                              120

      Leu
      121

25     <210> 99
      <211> 121
      <212> PRT
      <213> artificial sequence

30     <220>
      <221> artificial
      <222> 1-121
      <223> humanized antibody heavy chain variable domain

35     <400> 99
      Glu Val Gln Leu Val Glu Ser Gly Gly Gly Leu Val Gln Pro Gly
      1           5           10           15

40     Gly Ser Leu Arg Leu Ser Cys Ala Ala Ser Gly Tyr Asp Phe Thr
      20           25           30

      His Tyr Gly Met Asn Trp Val Arg Gln Ala Pro Gly Lys Gly Leu
45     35           40           45

      Glu Trp Val Gly Trp Ile Asn Thr Tyr Thr Gly Glu Pro Thr Tyr
      50           55           60

      Ala Ala Asp Phe Lys Arg Arg Phe Thr Phe Ser Leu Asp Thr Ser
50     65           70           75

      Lys Ser Thr Ala Tyr Leu Gln Met Asn Ser Leu Arg Ala Glu Asp
      80           85           90

55     Thr Ala Val Tyr Tyr Cys Ala Lys Tyr Pro His Tyr Tyr Val Asn
      95           100          105

      Glu Arg Lys Ser His Trp Tyr Phe Asp Val Trp Gly Gln Gly Thr
      110          115          120

```

WO 00/29584

PCT/US99/27153

Leu
121

5

INTERNATIONAL SEARCH REPORT

International Application No

PCT/US 99/27153

A. CLASSIFICATION OF SUBJECT MATTER

IPC 7 C12N15/13 C12N15/63 C07K16/00 A61K39/395 C12N5/10

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

IPC 7 C07K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	<p>WILSON PATRICK C ET AL: "Somatic hypermutation introduces insertions and deletions into immunoglobulin V genes." JOURNAL OF EXPERIMENTAL MEDICINE JAN. 5, 1998, vol. 187, no. 1, 5 January 1998 (1998-01-05), pages 59-70, XP002134538 ISSN: 0022-1007 the whole document</p> <p style="text-align: center;">--- -/-</p>	1-15, 18, 20-31

☒ Further documents are listed in the continuation of box C.

☒ Patent family members are listed in annex.

* Special categories of cited documents:

- *A* document defining the general state of the art which is not considered to be of particular relevance
- *E* earlier document but published on or after the international filing date
- *L* document which may throw doubt on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
- *O* document referring to an oral disclosure, use, exhibition or other means
- *P* document published prior to the international filing date but later than the priority date claimed

- *I* later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
- *X* document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
- *Y* document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.
- *G* document member of the same patent family

Date of the actual completion of the international search

31 March 2000

Date of mailing of the international search report

27/04/2000

Name and mailing address of the ISA

European Patent Office, P.B. 5818 Patentlaan 2
NL - 2280 HV Rijswijk
Tel. (+31-70) 340-2040, Tx. 31 651 epo nl.
Fax: (+31-70) 340-3016

Authorized officer

Covone, M

1

INTERNATIONAL SEARCH REPORT

Internat Application No
PCT/US 99/27153

C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	FEENEY A J ET AL: "Sequence and fine specificity analysis of primary 511 anti-phosphorylcholine antibodies." JOURNAL OF IMMUNOLOGY, (1989 DEC 15) 143 (12) 4061-8. , XP002134539 abstract page 4064, left-hand column, paragraph 2 -page 4065, left-hand column, paragraph 1 page 4067, left-hand column, line 55-61 ----	1-15, 18, 26-31
A	OHLIN MATS ET AL: "Insertions and deletions in hypervariable loops of antibody heavy chains contribute to molecular diversity." MOLECULAR IMMUNOLOGY MARCH, 1998, vol. 35, no. 4, March 1998 (1998-03), pages 233-238, XP000891714 ISSN: 0161-5890 page 233, right-hand column, line 23 -page 234, left-hand column, line 3 page 236, left-hand column, paragraph 2 -page 237, left-hand column, paragraph 3 figure 2 ----	1-31
A	PATENT ABSTRACTS OF JAPAN vol. 1997, no. 10, 31 October 1997 (1997-10-31) & JP 09 140386 A (EIKEN CHEM CO LTD), 3 June 1997 (1997-06-03) abstract -& DATABASE WPI Oerwent Publications Ltd., London, GB; XP002134540 & JP 09 140386 A abstract ----	1-31
A	CA 2 125 240 A (CANADA NAT RES COUNCIL) 7 December 1995 (1995-12-07) page 3, line 5-15 examples 1,2 -----	1-31

INTERNATIONAL SEARCH REPORT

Information on patent family members

International Application No
PCT/US 99/27153

Patent document cited in search report	Publication date	Patent family member(s)	Publication date
JP 09140386 A	03-06-1997	NONE	
CA 2125240 A	07-12-1995	NONE	